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Perfect score: Sequence:

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Run on:

Scoring table:

Searched:

Database

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Sequence 1, A Sequence 16, Sequence 16, Sequence 11, A Sequence 11, A
                   Sequence Sequence S
                                                Sequence Sequence 2
                                                                           Sequence 4
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Sequence
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Sequence
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                                                                                                                                                                                                                                                                                                                                                                                                GENERAL INFORMATION:

APPLICANT: WILLICK, Gordon E.

APPLICANT: WHITFIELD, James F.

APPLICANT: SUREWICZ, Witchd

APPLICANT: SUNG, WING L.

APPLICANT: NEUGENBAUER, Witchd

APPLICANT: NEUGENBAUER, Witchd

APPLICANT: PREATHYROID HORMONE ANALOGUES

TITLE OF INVENTION: FOR THE TREATMENT OF OSTEOPOROSIS

NUMBER OF SEQUENCES: 6

CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 134; DB 1;
Pred. No. 7.6e-13;
); Mismatches 1;
US-08-904-760B-22
US-08-903-497A-1
PCT-US95-15800-22
US-08-1256-363-4
US-08-112-024-2
US-08-126-363-4
US-08-440-117-1
US-08-08-738A-16
US-08-123-849-1
US-08-625-586-1
US-08-625-586-1
US-09-625-586-1
US-09-625-586-1
US-09-625-586-1
US-07-773-098-2
                                                                                                                                                                                                                       US-08-332-453-2
US-08-689-190-2
US-08-142-551B-1
                                                                                                                                                                                                           US-07-863-014-2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       COUNTRY: Canada
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
COMPUTER: IBM PC COMPATIBLE
SOFTWARE: Wordperfect 5.1
CURRENT APPLICATION NATA:
APPLICATION NUMBER: US/08/262,495C
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADDRESSEE: Kirby, Eades, Gale, Baker
STREET: 112 Kent Street, Suite 770,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 FILING DATE:
CLASSIFICATION:
ATTORNEY/AGENT INFORMATION:
NAME: EADES, NO. 5556940Tis M.
REGISTRATION NUMBER: 5,263
REFERENCE/DOCKET NUMBER: 36210
TELECOMMUNICATION INFORMATION:
TELEPHONE: (613)-237-6900
TELEPHONE: (613)-237-6005
INFORMATION FOR SEQ ID NO: 5:
                                                                                                                                                                                                                                                                                                                                                                       ; Sequence 5, Application US/08262495C
; Patent No. 5556940
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LENGTH: 30 amino acids
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 97.8
Best Local Similarity 96.3
Matches 26; Conservative
      PRIOR APPLICATION DATA: APPLICATION NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MOLECULE TYPE: peptide US-08-262-495C-5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           FILING DATE:
CLASSIFICATION: 530
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CITY: Ottawa
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TOPOLOGY:
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47.295 Million cell updates/sec
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 6, Ap
Sequence 14, A
Sequence 32, A
Sequence 1, Ap
Sequence 1, Ap
Sequence 1, Ap
                                                                                                         ; Search time 12.19 Seconds
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Sequence 6
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Sequence 2
Sequence 7
Sequence 6
Sequence 1
Sequence 2
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Sequence 1
Sequence 1
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Sequence 5
Sequence 1
Sequence 1
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/cgn2_6/ptodata/2/jaa/5B_COMB.pep:*
/cgn2_6/ptodata/2/jaa/6A_COMB.pep:*
/cgn2_6/ptodata/2/jaa/6B_COMB.pep:*
/cgn2_6/ptodata/2/jaa/PcTUS_COMB.pep:*
/cgn2_6/ptodata/2/jaa/PcTUS_COMB.pep:*
                   GenCore version 4.5
Copyright (c) 1993 - 2000 Compugen Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             US-08-262-495C-5

US-08-262-495C-3

US-08-691-647C-1

US-08-694-760B-1

US-08-904-760B-6

US-08-904-760B-1

US-08-904-760B-14

US-08-904-760B-14

US-08-904-765B-14

US-08-14-765-373-1

US-08-14-765-373-1

US-08-14-863-1

US-08-14-863-1

US-08-14-07-1

US-08-14-07-1
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-08-521-097-1
-09-044-536A-1
                                                                                                                                                               US-09-448-867-1
137
1 XVSEIQLMHNIGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                   197339 seqs, 20590346 residues
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                                                                                                                                                                                                                                                                                                                                                                  Post-processing: Minimum Match 08
Maximum Match 1008
Listing first 45 summaries
                                                                            protein search, using sw model
                                                                                                        August 31, 2001, 15:59:56
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Gapop 10.0 , Gapext 0.5
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Maximum DB seq length: 200000000
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Match
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Result

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APPLICANT: Neugebauer, Witold
APPLICANT: Ross, Virginia
APPLICANT: Whitfield, James
APPLICANT: Willick, Gordon E.
TITLE OF INVENTION: CYCLIC PARATHYROID HORMONE ANALOGUES
TITLE OF INVENTION: FOR THE TREATMENT OF OSTEOPOROSIS
CYCLIC PARATHYROID HORMONE ANALOGUES FOR THE TREATMENT OF OSTEOPOROSIS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Indels
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96.3%; Pred. No. 7.9e-13;
live 0; Mismatches 1;
                                                              CORRESPONDENCE ADDRESS:
ADDRESSEE: NIXON & VANDERHYE, P.C.
STREET: 1100 New York Avenue, 8th Floor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  E: NIXON & VANDERHYE, P.C.
1100 New York Avenue, 8th Floor
                                                                                                                                                                                                                                                                                                      SOFTWARE ASCII Text
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/691,647C
FILING DATE: August 2, 1996
CLASSIFICATION: 514
ATTORNEY/AGENT INFORMATION:
NAME: CTAWFORD, ATTORNEY/AGENT UNMBER: 25,327
REFERENCE/DOCKET NUMBER: 1339-5
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CURRENT APPLICATION DATA: APPLICATION NUMBER: US/08/691,647C
                                                                                                                                                      STATE: Virginia
COUNTRY: U.S.A.
ZIP. 22201-4714
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Sequence 6, Application US/08691647C Patent No. 5955425
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    APPLICANT: Barbier, Jean-Rene
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TELEPHONE: (703) 816-4005
TELEFAX: (703) 816-4100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Floppy disk
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Best Local Similarity 96.3
Matches 26; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CITY: Arlington
STATE: Virginia
COUNTRY: U.S.A.
ZIP: 22201-4714
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy di
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            INFORMATION FOR SEQ ID NO:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQUENCE CHARACTERISTICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ASCII Text
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CORRESPONDENCE ADDRESS:
                                               NUMBER OF SEQUENCES:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        linear
                                                                                                                                    CITY: Arlington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADDRESSEE:
STREET: 11
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TOPOLOGY:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TELEFAX:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SOFTWARE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       LENGTH:
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Pred. No. 7.9e-13;
0; Mismatches 1; Indels
                                                                                                                                                                                                              GENERAL INFORMATION:
APPLICAMY: WILLICK, Gordon E.
APPLICAMY: WHITFIELD, James F.
APPLICAMY: SUREWICZ, Witcold
APPLICAMY: SUNG, Wing L.
APPLICAMY: NEUGENBAUER, Witcold
TITLE OF INVENTION: FOR THE TREATMENT OF OSTEOPOROSIS
                                                                                                                                                                                                                                                                                                                                                                                                                                        ADDRESSEE: Kirby, Eades, Gale, Baker
STREET: 112 Kent Street, Suite 770,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SCETWARE: Wordperfect 5.1
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/262,495C
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         CLASSIFICATION: 530
ATTORNEY/AGENT INFORMATION:
NAME: EADES, NO. 5556940ris M.
REGISTRATION NUMBER: 5,263
REFERENCE/DOCKET NUMBER: 36210
TELEPHONE: (613)-237-6900
TELEPHONE: (613)-237-6900
TELEPHONE: (613)-237-0045
: SEQUENTION FOR SEQ ID NO: 3:
SEQUENTION FOR SEQ ID NO: 3:
SEQUENTION FOR SEQ ID NO: 3:
                                            2 VSEIQLMHNLGKHLNSMERVEWLRKKL 28
                        2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  COUNTRY: Canada
COMPUTER REAABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC Compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Sequence 1, Application US/08691647C
Patent No. 595442
GENERAL INFORMATION:
APPLICANT: Barbier, Jean-Rene
                                                                                                                                                                        ; Sequence 3, Application US/08262495C
; Patent No. 5556940
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Neugebauer, Witold
Ross, Virginia
Whitfield, James
Willick, Gordon E.
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Best Local Similarity 96.3%;
Matches 26; Conservative (
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CLASSIFICATION: 530
PRIOR APPLICATION DATA:
APPLICATION NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TOPOLOGY: linear
MOLECULE TYPE: peptide
                                                                                                                                                                                                                                                                                                                                                                                           NUMBER OF SEQUENCES: 6
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Ottawa
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  FILING DATE:
                                                                                                                                               US-08-262-495C-3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               US-08-262-495C-3
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US-08-691-647C-1
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APPLICANT:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   APPLICANT:
APPLICANT:
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RESULT

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Gaps

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APPLICANT: Jean-Rene, Barbier
APPLICANT: Neugebauer, Witold
APPLICANT: Ross, Virginia
APPLICANT: Whitfield, James
APPLICANT: Whitfield, James
APPLICANT: WILLICK, Gordon E.
TITLE OF INVENTION: PARATHYROID HORMONE ANALOGUES FOR THE
TITLE OF INVENTION: TREATMENT OF OSTEOPOROSIS
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        cyclo Lys27-Asp30, and this sequence has an amino group c-terminus (NH2).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           97.8%; Score 134; DB 3; Length 31; 96.3%; Pred. No. 7.9e-13; Live 0; Mismatches 1; Indels
                                                                                Score 134; DB 3; Length 31;
Pred. No. 7.9e-13;
0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADDRESSEE: NIXON & VANDERHYE P.C.
STREET: 1100 No. 6110892th Glebe Rd. 8th floor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
COMPATION SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
FILING DATE: 01-AUG-1997
CLASSIFICATION: 514
                                                                                                                                                                                            2 VSEIQLMHNLGKHLNSMERVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          2 VSEIQLMHNLGKHLNSMERVEWLRKKL 28
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                                                                                                                                                                                                                                                                                                                 Sequence 6, Application US/08904760B Patent No. 6110892 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/691,647
FILING DATE: 02-AUG-1996
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            REGISTRATION NUMBER: 25,327
REFERENCE/DOCKET NUMBER: 13
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       TELECOMMUNICATION INFORMATION
                                                                                  97.8%;
96.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        NAME: Crawford, Arthur R
                                                                         Ouery Match
Best Local Similarity 96.39
warrhes 26; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TELEPHONE: 703-816-400
TELEFAX: 703-816-4100
INFORMATION FOR SEQ ID NO:
; MOLECULE TYPE: protein US-08-904-7608-1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ZIP: 22201-4741
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             circular
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OTHER INFORMATION:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               STREET: 1100 No. CITY: Arlington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
Matches 26; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               FILING DATE: 01 CLASSIFICATION:
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APPLICANT: Jean-Rene, Barbier
APPLICANT: Neugebauer, Witcld
APPLICANT: Ross, Virginia
APPLICANT: Whitfield, James
APPLICANT: Willick, Gordon E.
TITLE OF INVENTION: PARATHYROID HORMONE ANALOGUES FOR THE
TITLE OF INVENTION: TREATMENT OF OSTEOPOROSIS
                                                                                                                                                                                                                                                                                                                                                                                                     Score 134; DB 2; Length 31;
Pred. No. 7.9e-13;
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COUNTRY: USA
ZIP: 22201-4741
ZIP: Z2201-4741
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION NUMBER: US/08/904,760B
TLING DATE: 01-AUG-1997
TLING DATE: 01-AUG-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                              0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        E: NIXON & VANDERHYE P.C.
1100 No. 6110892th Glebe Rd.
                                      ATTORNEY/AGENT INFORMATION:
NAME: Crawford, Arthur R.
REGISSRATION NUMBER: 25,327
REFERENCE/DOCKET NUMBER: 1339-5
TELECOMMUNICATION INFORMATION:
TELEPHONE: (703) 816-4005
MILLEPLECAX: (703) 816-4100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2 VSEIQLMHNIGKHLNSMERVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PRIOR APPLICATION DATA:
APPLICATION DATA:
APPLICATION NUMBER: 08/691,647
FILING DATE: 02-AUG-1996
ATTORNEY AGENT INFORMATION:
NAME: Crawford, Arthur R.
REGISTRATION NUMBER: 25,327
REFERENCE/DOCKET NUMBER: 1339-6
TELECOMMUNICATION INFORMATION:
TELECHONE: 703-816-4000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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August 2, 1996
N: 514
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INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                   SEQUENCE CHARACTERISTICS:
LENGTH: 31 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                            26; Conservative
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                                                                                                                                                                                       TELEX: N/A INFORMATION FOR SEQ ID NO:
                                                                                                                                                                                                                                                                                             TOPOLOGY: cyclic
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NUMBER OF SEQUENCES: 3
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                            amino acid
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Best Local Similarity
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  FILING DATE: A
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADDRESSEE:
                                                                                                                                                                                                                                                                                                                                      US-08-691-647C-6
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Gaps

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97.8%; Score 134; DB 3; Length 31; 96.3%; Pred. No. 7.9e-13;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                This sequence has an amino group c-terminus (NH2).
                                                                                                                                                                                                        MEDIUM TYPE: Floppy disk COMBUTER: BM PC compatible COMBTER: PC-DOS/MS-DOS SOFTWARE: Patentin Release #1.0, Version #1.30 CURRENT APPLICATION DATA: APPLICATION NUMBER: US/08/904,760B FILING DATE: 01-AUG-1997 CLASSIFICATION: 51 PRIOR APPLICATION DATA: APPLICATION DATA: APPLICATION NUMBER: 08/691,647
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APPLICANT: FUKUDA, Tsunehiko
APPLICANT: KAWASE, Masahiro
APPLICANT: YAMAZAKI, IWAO
APPLICANT: YAMAZAKI, IWAO
TITLE OF INVENTION: PARATHYROID HORMONE DERIVATIVES
NUMBER OF SEQUENCES: 2
                                                                                   8th floor
TITLE OF INVENTION: TREATMENT OF OSTEOPOROSIS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               0; Mismatches
                                                             E: NIXON & VANDERHYE P.C.
1100 No. 6110892th Glebe Rd.
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                                                                                                                                                                                                                                                                                                                                                                                                                  FILING DATE: 02-AUG-1996
ATTORNEY/AGENT INFORMATION:
NAME: Crawford, Arthur R.
REGISTRATION NUMBER: 25,327
REFERENCE/DOCKET NUMBER: 1339-6
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Patent No. 5393869
GENERAL INFORMATION:
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TELECOMMUNICATION INFORMATION:
TELEPHONE: 703-816-4000
TELEFAX: 703-816-4100
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 SEQUENCE CHARACTERISTICS:
LENGTH: 31 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Conservative
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                                                                                                                                            COUNTRY: USA
ZIP: 22201-4741
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TOPOLOGY: linear
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CITY: Boston
STATE: Massachusetts
                   NUMBER OF SEQUENCES: 3 CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CORRESPONDENCE ADDRESS:
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; OTHER INFORMATION:
US-08-904-7608-32
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best_Local Similarity
Matches 26; Conserv
                                                                                                       Arlington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             STRANDEDNESS
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                                                               ADDRESSEE:
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                                                  Sequence 14, Application US/08904760B
Patent No. 6110892
GENERAL INFORMATION:
APPLICANT: Jean-Rene, Barbier
APPLICANT: Magebauer, Witcold
APPLICANT: Whitfield, James
APPLICANT: Whitfield, James
APPLICANT: Whitlick, Gordon E.
TITLE OF INVENTION: PARATHYROID HORMONE ANALOGUES FOR THE
TITLE OF INVENTION: TREATMENT OF OSTEOPOROSIS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              APPLICANT: Neugebauer, Witold
APPLICANT: Ross, Virginia
APPLICANT: Whitfield, James
APPLICANT: Whilfick, Gordon E.
TITLE OF INVENTION: PARATHYROID HORMONE ANALOGUES FOR THE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       cyclo Glu22-Lys26, and this sequence has an amino group c-terminus (NH2).
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Pred. No. 7.9e-13;
0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                          STALL.
COUNTRY: USA
ZIP: 22201-4741
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBAM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
STELIGATION UNDER: US/08/904,760B
TING DATE: 01-AUG-1997
                                                                                                                                                                                                                                                                                                                                       8th floor
                                                                                                                                                                                                                                                                                                                 ADDRESSEE: NIXON & VANDERHYE P.C.
STREET: 1100 No. 6110892th Glebe Rd.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PRIOR APPLICATION DATA:
PRPLICATION NUMBER: 08/691,647
FILING DATE: 02-AGG-1996
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    NAME: Crawford, Arthur R. REGISTRATION NUMBER: 25,327 REFERENCE/DOCKET NUMBER: 135 TELECOMMUNICATION INFORMATION: TELEPHONE: 703-816-4000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            APPLICANT: Jean-Rene, Barbier
APPLICANT: Neugebauer, Witold
APPLICANT: Ross, Virginia
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             INFORMATION FOR SEQ ID NO: 14: SEQUENCE CHARACTERISTICS:
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96.3%;
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amino acid
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Best Local Similarity 96.3'
Matches 26; Conservative
                                                                                                                                                                                                                                                                       NUMBER OF SEQUENCES: 35
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           TELEPHONE: 703-516-4100
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TOPOLOGY: circular
MOLECULE TYPE: protein
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; OTHER INFORMATION:
US-08-904-760B-14
                                                                                                                                                                                                                                                                                                                                                           Arlington
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US-08-904-760B-32
               RESULT 7
US-08-904-760B-14
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                                                                                     Score 134; DB 1; Length 34;
Pred. No. 8.7e-13;
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                                                                                                                                                                                                                                                                                                                                                                                                                                         APPLICANT: SUNG, WING L.
APPLICANT: NEUGENBAUER, WILCOLD
TITLE OF INVENTION: PARATHYROID HORMONE ANALOGUES
TITLE OF INVENTION: FOR THE TREATMENT OF OSTEOPOROSIS
NUMBER OF SEQUENCES: 6
CORRESPONDENCE ADDRESS:
                                                                                                                              0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            COMPRY: Canada
COMPURER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Wordperfect 5.1
CURRENT APPLICATION NATA:
APPLICATION NUMBER: US/08/262,495C
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       E: Kirby, Eades, Gale, Baker
112 Kent Street, Suite 770,
                                                                                                                                                                                           2 VSEIQLMHNLGKHLNSMERVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        2 VSEIQLMHNLGKHLNSMERVEWLRKKL 28
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      REFERENCE/DOCKET NUMBER: 36210
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                                                                                                                                                                                                                                                                                                                   ; Sequence 1, Application US/08262495C
; Patent No. 5556940
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ATTORNEY/AGENT INFORMATION:
NAME: EADES, NO. 5556940ris M.
REGISTRATION NUMBER: 5,263
                                                                                                                                                                                                                                                                                                                                                          GENERAL INFORMATION:
APPLICANT: WILLICK, GOTGOD E.
APPLICANT: WHITFIELD, James F.
APPLICANT: SUREWICZ, Witold
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TELECOMMUNICATION INFORMATION:
                                                                                   97.88;
96.38;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (613)-237-6900
(613)-237-0045
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             97.8%;
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; FRAGMENT TYPE: N-terminal US-08-033-099-1
                                                                                                        Best Local Similarity 96.3
Matches 26; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SEQUENCE CHARACTERISTICS:
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Best Local Similarity 96.3
Matches 26; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           PRIOR APPLICATION DATA: APPLICATION NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TOPOLOGY: linear
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          CLASSIFICATION: 530
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         CLASSIFICATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Ottawa
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    FILING DATE:
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                                                                                                                                                                                                                                                                           RESULT 11
US-08-262-495C-1
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                                                                                       Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 APPLICANT: NAKAGAWA, Shizue
APPLICANT: TAKETOMI, Shigehisa
TITLE OF INVENTION: PARATHYROID HORMONE DERIVATIVES
NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
ADDRESSEE: DAVID G.CONLIN; DIKE, BRONSTEIN, ROBERTS
ADDRESSEE: CUSHWAN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: TBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/033,099
FILING DATE: 19930316
CLASSIFICATION: 530
  SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/07/765,373
FILING DATE: 19910925
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                                                                                                    ATTORNEY/AGENT INPORMATION:
NAME: WILLIAMS, Gregory D.
REGISTRATION UNDRER: 30901
REFERENCE/DOCKET NUMBER: 41289
TELECOMMUNICATION INFORMATION:
TELEPHONE: (617)523-3400
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     REFERENCE/DOCKET NUMBER: 42528 TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 1, Application US/08033099
Patent No. 5434246
GENERAL INFORMATION:
APPLICANT: FUKUDA, Tsunehiko
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ATTORNEY AGENT INFORMATION:
NAME: WILLIAMS, Gregory D
REGISTRATION NUMBER: 30901
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     97.8%;
96.3%;
                                                                                                                                                                                                                              TELEFAX: (617)523-6440
- TELEX: 20091 STRE UR
INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            TELEPHONE: (617)523-3400
TELEFAX: (613)523-6440
TELEY: 200291 STRE UR
INFORMATION FOR SEQ ID NO: 1:
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                                                                                                                                                                                                                                                                                                                                                                             peptide
N-terminal
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Best Local Similarity 96.3
Matches 26; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Massachusetts
                                                                                                                                                                                                                                                                                                LENGTH: 34 and Toner: AMINO ACID
                                                             FILING DATE: 19
CLASSIFICATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Boston
                                                                                                                                                                                                                                                                                                                                                                             MOLECULE TYPE:
FRAGMENT TYPE:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              US-08-033-099-1
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
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                                                                                                                                                                                                                                                                                                                                                                                                     amino acid
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HYPOTHETICAL:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ; FRAGMENT TYPE:
US-08-443-863-1
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CORRESPONDENCE: 34

CORRESPONDENCE ADDRESS:
                                                                                                                TITLE OF INVENTION: ANALOGS OF PARATHYROID HORMONE AND TITLE OF INVENTION: PARATHYROID HORMONE RELATED PEPTIDE: SYNTHESIS AND USE TITLE OF INVENTION: FOR THE TREATMENT OF OSTEOPOROSIS

WUMBER OF SEQUENCES: 34
                                                                                                                                                                                                                                                                                                                                                                                                                 SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/07/915,247A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Patent Dept., Syntex (U.S.A.), Inc
                                                                                                                                                                                                                      Patent Dept., Syntex (U.S.A.), Inc
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Pred. No. 8
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19920714
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US-08-443-863-1
; Sequence 1, Application US/08443863
; Patent No. 5693616
; GENÉRAL INFORMATION:
; APPLICANT: Krstenansky, John L.
APPLICANT: Nestor Jr., John J.
APPLICANT: Ho, Teresa H.
; APPLICANT: Wickery, Brian H.
; APPLICANT: Wickery, Brian H.
                                                                                                                                                                                                                                                                                                                                                           MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
OPERATING SYSTEM: PC-DOS/MS-DOS
                      Krstenansky, John L.
                                   APPLICANT: Nestor Jr., John J.
APPLICANT: HO, Teresa H.
VICKERY, BIAN H.
APPLICANT: Bach, Chinh T.
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TELECOMMUNICATION INFORMATION:
TELEPHONE: 415-855-6593
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96.3%;
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Palo Alto
                                                                                                                                                                                                                                       STREET: 3401 Hillview Ave.
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NAME: Schmonsees, William
REGISTRATION NUMBER: 31,75
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           LENGTH: 34 amino acids
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 97.8
Best Local Similarity 96.3
Matches 26; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            TELEFAX: 415-496-3529
INFORMATION FOR SEQ ID NO:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                      COMPUTER READABLE FORM: MEDIUM TYPE: Floppy
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   MOLECULE TYPE: peptide
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                linear
GENERAL INFORMATION:
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                                                                                                                                                                                                                                                                                                   USA
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FRAGMENT TYPE:
                                                                                                                                                                                                                                                                                                                    94303
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ANALOGS OF PARATHYROID HORMONE AND PARATHYROID HORMONE RELATED PEPTIDE: SYNTHESIS AND USE FOR THE TREATMENT OF OSTEOPOROSIS
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                                                 SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/443,863
FILING DATE: 14-JUL-1992
CLASSIFICATION: 514
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CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/448,070
FILING DATE: U4-JUL-1992
CLASSIFICATION: 435
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               97.8%; Score 134; 96.3%; Pred. No. 8
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COMPUTER: IBM PC compatible OPERATING SYSTEM: PC-DOS/MS-DOS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 1, Application US/08448070
Patent No. 5655955
GENERAL INFORMATION:
APPLICANT: Krstenansky, John L.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          APPLICANT: Nestor Jr., John J.
APPLICANT: Ho, Teresa H.
APPLICANT: Vickery, Brian H.
APPLICANT: Bach, Chinh T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       REGISTRATION NUMBER: 31,796
REFERENCE/DOCKET NUMBER: 276
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                                                            NAME: Schmonsees, William REGISTRATION NUMBER: 31,796 REFERENCE/POCKET NUMBER: 276 TELECOMMUNICATION INFORMATION: TELEPHONE: 415-855-5593 TELEFAX: 415-496-3529
                                                                                                                                                                                                                                                                                                                                             TELEFAX: 415-496-3529
INFORMATION FOR SEQ ID NO: 1:
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                                                                                                                                                                                                                                                                  97.8%; Score 134; DB 1; Length 34; 96.3%; Pred. No. 8.7e-13; Live 0; Mismatches 1; Indels
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Pred. No. 8.7e-13;
0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
COMPUTER: IBM PC COMPATIBLE
COMPUTER: IBM PC COMPATIBLE
COMPUTER: IBM PC COMPATIBLE
COMPATING SYSTEM: PC-POS/NS-DOS
SOFTWARE: PATENTIN Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/488,105
FILING DATE: US/09/488,105
FILING DATE: US/09/488,105
FILING DATE: US/09/488,105
FILING DATE: US/08/488,105
FILING DATE: US/08/488,105
TELESERYION NUMBER: 34,053
REGISTRATION NUMBER: 34,053
REGISTRATION INFORMATION:
TELEPRAK: 617/542-8906
TELEPRAK: 617/542-8906
TELEPRAK: 617/542-8906
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 7, Application US/08488105
Fatent No. 5717062
GENERAL INFORMATION:
APPLICANT: Chorev, Michael
TITLE OF INVENTION: CYCLIC ANALOGS OF PTH AND PTHIP
NUMBER OF SEQUENCES: 22
CORRESPONDENCE ADDRESS:
ADDRESSEE: Fish & Richardson P.C.
STREET: 225 Franklin Street
                                                                                                                                                                                                                                                                                                                                                           2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                  2 VSEIQLMHNIGKHLNSMERVEWLRKKL 28
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TELEFAX: 415-496-3529 INFORMATION FOR SEQ ID NO: 1:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            97.8%;
96.3%;
                                                                                                                                                                                N-terminal
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TELEX: 200154
INFORMATION FOR SEQ ID NO:
SEQUENCE CHARACTERISTICS:
LENGTH: 34 aming acids
                                         SEQUENCE CHARACTERISTICS:
LENGTH: 34 amino acids
                                                                                                                                                                                                                                                                                         Best_Local Similarity 96.3
Matches 26; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match 97.8
*Best Local Similarity 96.3
Matches 26; Conservative
                                                                                  TYPE: amino acid
TOPOLOGY: linear
MOLECULE TYPE: peptide
HYPOTHETICAL: NO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  COUNTRY: USA
ZIP: 02110-2804
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy (
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MOLECULE TYPE: protein
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US-08-488-105-7
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STRANDEDNESS: no
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US-08-448-070-1
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US-08-488-105-7
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Search completed: August 31, 2001, 16:00:17 Job time: 21 sec

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Perfect score:

Sequence:

OM protein -

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Scoring table:

Searched:

Minimum DB Maximum DB

Database

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Human parathyroid hormone; hPTH; variant; analogue;
calcium; depletion; fixation; resorption; osteopathy; osteoporosis;
hypoparathyroidism.
                                                                                                                                                                                                                                                                                                                                                                                           ALIGNMENTS
                                                                                                                             AAR07919
AAR07922
AAR1248
AAR1549
AAR58291
AAR58291
AAR5828
AAR5828
AAR58181
AAR58016
AAR58016
AAR58016
                                                    AAY 02578
AAY 98012
AAY 98015
AAY 98018
AAP 30022
AAP 50377
AAP 60031
                                                                                                                                                                                                                                                            AAR74521
AAW99449
AAR99978
AAR98951
AAR98966
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AAW20000
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       92GB-0015009.
92GB-0026415.
92GB-0026859.
92GB-0001691.
93GB-0001692.
93GB-0007673.
93GB-0014384
[Ala19]-hPTH(1-36)-NH2
Key
Modified-site
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       23-DEC-1992;
28-JAN-1993;
28-JAN-1993;
14-APR-1993;
19-APR-1993;
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23-DEC-1992
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Synthetic.
AAR58214;
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 RESULT
                                                                                                                                                                                                                                                                                                                                                                                                                                                [Ala19]-hPTH(1-36)
[Ser19]-hPTH(1-38)
[Ala19]-hPTH(1-38)
Human parathyroid
Human parathyroid
Human parathyroid
Human parathyroid
Human parathyroid
Human parathyroid
                                                                       Search time 18.06 Seconds (without alignments) 93.991 Million cell updates/sec
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Copyright (c) 1993 - 2000 Compugen Ltd.
                                                                                                                                        1 XVSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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                                                                                                                                                                                              412676 seqs, 60623988 residues
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Maximum Match 100%
Listing first 45 summaries
                                                      protein search, using sw model
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AAY98042
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Gapop 10.0, Gapext 0.5
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                                                                          August 31,
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Match
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135 135 134 134 134 134 134 134

Score

Result Š

[Lys(For)26, Lys(F [D-Asp30]-hPTH(1-3 [Lys32]-hPTH(1-34) [Thr33, Ala34]-hPT

Parathyroid hormon [D-Ser3]hPTH (1-34 [Gln25]hPTH (1-34)

Human parathyroid

Human parathyroid-[Met(0)8,18]hPTH-(Sequence of the fi Human parathyroid

Human amino-termin amino-termin Human amino-termin

N-terminal 31 resi

Human parathyroid Human parathyroid Human parathyroid

Adenine-rich PTH-

Human parathyroid Human parathyroid Target peptide (PT PTH(1-34). Not sp

PTH(1-34). Not sp Human parathyroid

parathyr

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Parathyroid hormon

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Human parathyroid

N-alpha-Isopropyl-

Gaps

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Human parathyroid hormone; hPTH; variant; analogue;
calcium; depletion; fixation; resorption; osteopathy; osteoporosis;
                                                                                                                                           This peptide is an example of a highly generic formula covering parathyroid hormone variants useful for treating or preventing bone conditions associated with calcium depletion/resorption, in cases where calcium fixation is required (esp. osteoporosis) or to treat hypoparathyroidism.
                                                               New active para-thyroid hormone variants - used for treating or preventing osteoporosis etc.
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                                                                                                                                                                                                                                                                                                  Length 38;
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Lewis I, Ramage P, Schneider
                                                                                                                                                                                                                                                                                              Score 135; DB 15;
Pred. No. 3.7e-13;
); Mismatches 1;
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SANDOZ PATENT GMBH.
SANDOZ-ERFINDUNGEN VERW GES MBH.
                                                                                                             Example 120; Page 39; 92pp; English.
                                                                                                                                                                                                                                                                                                                                                               2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                2 vseiglmhnlqkhlnsmsrvewlrki 28
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92GB-0026859.
92GB-0026861.
93GB-0001691.
93GB-0007673.
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96.3%;
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Best Local Similarity 96.39
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Gombert F, Gram H, Le
Waelchli R, Rainer A;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 [Ala19]-hPTH(1-38)-OH.
                                WPI; 1994-018352/03
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                hypoparathyroidism
                                                                                                                                                                                                                                                38 AA;
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14-APR-1993;
Waelchli R,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human parathyroid hormone; hPTH; variant; analogue;
calcium; depletion; fixation; resorption; osteopathy; osteoporosis;
                                                                                                                                                                                            New active para-thyroid hormone variants - used for treating or preventing osteoporosis etc.
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0
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Lewis I, Ramage P, Schneider H;
                                                                                              lge.R, Cardinaux F;
Ramage P, Schneider
                                                                                                                                                                                                                                                                                                                                                                                                                                                               0; Mismatches
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SANDOZ-ERFINDUNGEN VERW GES MBH.
                           BAUER W.
SANDOZ PATENT GMBH.
SANDOZ-ERFINDUNGEN VERW GES MBH.
                                                                                            Breckenridge.R,
Lewis I, Ramage
                                                                                                                                                                                                                                            Example 212; Page 44; 92pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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93GB-0001691.
93GB-0001692.
93GB-0007673.
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                                                                                          Bauer W, Br
Gram H, Le
, Rainer A;
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Gram H,
                                                                                                                                                            WPI; 1994-018352/03
             SANDOZ LTD.
                                                                                                                                                                                                                                                                                                                                              hypoparathyroidism
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Matches 26; Conserv
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Gombert F, Gr
Waelchli R, F
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28-JAN-1993;
14-APR-1993;
19-APR-1993;
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18-DEC-1992;
23-DEC-1992;
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Gombert F,
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                        (SANO) S
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RESULT AAR58123

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osteopaenia; hypoparathyroidism; fracture repair; hypercalcaemia; breast cancer; lung cancer; prostate cancer; multiple myeloma; hypernephroma; had and neck epidermoid cancer; oesophagus cancer; osteopathic; PTH.
                                                                                                                                                                                                                                             Human; parathyroid hormone; signal transduction; osteoporosis;
                                                                                                                                                                                                              Human parathyroid hormone peptide # 1.
                         2 vseiglmhnlgkhlnsmervewlrkkl
         2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                      AAY98041 standard; peptide; 28
                                                                                                                                                                              04-SEP-2000 (first entry)
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                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     New human parathyroid hormone analogues - which have increased adenylyl cyclase activating activity, used for treating osteoporosis
This peptide is an example of a highly generic formula covering parathyroid hormone variants useful for treating or preventing bone conditions associated with calcium depletion/resorption, in cases where calcium fixation is required (esp. osteoporosis) or to treat hypoparathyroidism.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAR88829-R88841 are human parathyroid hormone (hPTH) analogues. The analogues increase G-protein coupled adenylyl cyclase (CAMPase) activity and reduce protein kinase C (PKC) activity. The analogues can reverse the loss of bone and increase bone mass and density without undesirable effects. They are useful for the treatment of osteoporosis and other bone related disorders and disorders
                                                                                                                                                                                   ö
                                                                                                                                                                                                                                                                                                                                                                                                                                              Parathyroid hormone; PTH; analogue; osteoporosis; bone cell; calcium regulation; reduced PKC activity; protein kinase C; increased adenylyl cyclase activity; cAMPase; bone loss.
                                                                                                                                                   Length 38;
                                                                                                                                                                                   Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Surewicz W, Whitfield JF;
                                                                                                                                                                                                                                                                                                                                                                                                                  Human parathyroid hormone analogue, hPTH(1-28)-NH2.
                                                                                                                                                 Score 135; DB 15;
Pred. No. 3.7e-13;
0; Mismatches 1;
                                                                                                                                                                                 0; Mismatches
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                                                                                                                                                                                                                                2 vseiglmhnlgkhlnsmarvewlrkkl
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                                                                                                                                                 98.5%;
96.3%;
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                                                                                                                                                                Local Similarity 96.3
nes 26; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          28 AA;
                                                                                                      38 AA;
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Modified-site
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Matches
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Potts

Gardella TJ,

G1y

label= Ser, Ala,

/label= Glu,

99WO-US27863

Location/Qualifiers

AA.

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carboxyl terminal of PTH is important for PTH receptor binding, while the carboxyl terminal of PTH is important for Signal transduction. It has been identified that the carboxyl terminal is important for Signal transduction. The present sequence is a human PTH peptide, with amino- and carboxy terminal captured to downstream signalling. Aberrant PTH activity has been implicated in a number of disorders: osteoporosis, osteopeania, hypoparathyroidism and hypercalcaemia. In turn, hypercalcaemia is associated with hyperrephroma and a variety of cancers: breast, lung and prostate carcinoma, multiple myeloma and epidermoid cancers of the head, neck and oesophagus. This peptide may be suitable for treatment of the fracture repair. Fragments of the present sequence would be suitable for fracture repair. Fragments of the present sequence are claimed e.g. fragments 1-24, 1-25, 1-26 and 1-27), providing that peptides are claimed e.g. or hPTH(1-26)NH2, hPTH(1-28)NH2.
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                                                                                                                                                                              Parathyroid hormone (PTH) binds to PTH receptors in renal and osseous
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Gaps
Novel biologically active peptide comprising a parathyroid hormone peptide derivative, useful for treating osteoporosis -
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Pred. No. 3.7e-13;
0; Mismatches 0;
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100.0%; Pre
                                                                                                           Claim 1; Page 67; 75pp; English.
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Best Local Similarity 100.
Matches 27; Conservative
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Gaps

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Length 28;

Score 134; DB 17; Length 2 Pred. No. 3.7e-13; 0; Mismatches 1; Indels

97.8%; 96.3%;

26; Conservative

Vouery Match Best Local Similarity Matches 26; Conserv

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cells, initiating signal transduction. It has been identified that the carboxyl terminal of PTH is important for PTH receptor binding, while the amino terminal is important for signal transduction. Various PTH peptides were produces with amino- and carboxy terminal modifications which had varying PTH receptor activation properties and therefore downstream signalling. Aberrant PTH activity has been implicated in a number of disorders: osteoporosis, osteopaenia, hypoparathyroidism and hypercalcaemia. In turn, hypercalcaemia is associated with hypernephroma and epidermoid cancers of the head, neck and oscophagus. The present sequence is a PTH peptide, with a Gly residue at position 1 and a Arg residue at position 19. The Gly residue reduces downstream is signalling via phospholipase C (PLC), whereas the Arg residue restores PLC signalling and ligand binding. The present peptide may therefore be used as a PTH receptor agonist for the treatment of the above mentioned disorders. In addition, the present sequence would be suitable for
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2 VSEIQLMHNLGKHLNSMXRVEWLRKKL
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                                                                                                                            AAY98042 standard; peptide; 28
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Novel biologically active peptide comprising a parathyroid hormone peptide derivative, useful for treating osteoporosis -

Claim 6; Page 68; 75pp; English.

Potts JT;

Gardella TJ,

Bringhurst FR, Takasu H, (GEHO) GEN HOSPITAL CORP.

WPI; 2000-400076/34.

99WO-US27863. 98US-0109938.

24-NOV-1999; 25-NOV-1998;

02-JUN-2000

osteopaenia; hypoparathyroidism; fracture repair; hypercalcaemia; breast cancer; lung cancer; prostate cancer; multiple myeloma; hypernephroma; head and neck epidermoid cancer; oesophagus cancer;

osteopathic; PTH.

WO200031266-A1.

Human; parathyroid hormone; signal transduction; osteoporosis;

Human parathyroid hormone peptide # 4.

(first entry)

04-SEP-2000

AAY98044;

AAY98044 standard; peptide; 28 AA

RESULT AAY98044

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cells, initiating signal transduction. It has been identified that the carboxyl terminal of PTH is important for PTH receptor binding, while the amino terminal is important for signal transduction. Various PTH peptides were produces with amino- and carboxy terminal modifications which had varying PTH receptor activation properties and therefore downstream is signalling. Aberrant PTH activity has been implicated in a number of disorders: osteoporosis, osteopaenia, hypoparathyroidism and hypercalcaemia. In turn, hypercalcaemia is associated with hypernephroma and a variety of cancers: breast, lung and prostate carcinoma, multiple myelome and epidermoid cancers of the head, neck and oscophagus. The present sequence is a PTH peptide, with an Ala residue at position 1 and an Arg residue at position 19. The Ala and Arg residues both improve constream signalling via phospholipase C (PLC) and ligand binding. The present peptide may therefore be used as a PTH receptor agonist for the treatment of the above mentioned disorders. In addition, the present sequence would be suitable for fracture repair.
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Gaps

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Length 28; Indels

97.8%; Score 134; DB 21; 96.3%; Pred. No. 3.7e-13; 0; Mismatches

Conservative

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breast cancer; lung
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                                             Human; parathyroid hormone; signal transduction; osteoporosis; osteopaenia; hypoparathyroidism; fracture repair; hypercalcaemia; breast cancer; lung cancer; prostate cancer; multiple myeloma; hypernephroma; head and neck epidermoid cancer; oesophagus cancer;
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Pred. No. 3.7e-13;
0; Mismatches 1;
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                       Human parathyroid hormone peptide # 6.
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04-SEP-2000 (first entry)
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                                                                                                                         Homo sapiens
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disorders: osteoporosis, osteopania, hypoparathyroidism and hypercalcaemia. In turn, hypercalcaemia is associated with hypernephroma and a variety of cancers: breast, lung and prostate carcinoma, multiple myeloma and epidermoid cancers of the head, neck and oesophagus. The present sequence is a PTH peptide, with an Ala residue at position 1 and a Glu residue at position 19. The Ala residue improved omstream signalling via phospholipase C (PLC), whereas the Glu residue reduces PLC signalling and ligand binding. PTH peptides with a Arg residue at position used as a PTH receptor agonist for the treatment of the above mentioned disorders and fracture repair.
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breast cancer; lung cancer; prostate cancer; multiple myeloma; hypernephroma; head and neck epidermoid cancer; oesophagus cancer;
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Pred. No. 3.7e-13;
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96.3%;
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Best Local Similarity 96.3
Matches 26; Conservative
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                                                                        osteopathic; PTH.
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                                                                                                                                                 Homo sapiens
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Gardella TJ,

(GEHO) GEN HOSPITAL CORP. Bringhurst FR, Takasu H,

WPI; 2000-400076/34

99WO-US27863 98US-0109938.

24-NOV-1999; 02-JUN-2000

WO200031266-A1

Claim 7; Page 69; 75pp; English

hypercalcaemia.

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Parathyroid hormone (PTH) binds to PTH receptors in renal and osseous cells, initiating signal transduction. It has been identified that the carboxyl terminal of PTH is important for PTH receptor binding, while the amino terminal is important for squal transduction. Various PTH peptides were produces with amino- and carboxy terminal modifications which had varying PTH receptor activation properties and therefore downstream signalling. Aberrant PTH activity has been implicated in a number of disorders: osteoporosis, osteopaenia, hypoparathyroidism and present and a variety of cancers: breast, lung and prostate carcinoma, multiple myeloma and epidermoid cancers of the head, neck and oesophagus. The myeloma and epidermoid cancers of the head, neck and oesophagus. The present sequence is a PTH peptide, with a Ser residue at position 1 and a Glu residue at position 19. The Ser residue improves downstream signalling and ligand binding. PTH peptides with a Arg residue at position part peptides with a Arg residue at position and ligand binding. PTH peptides with a Arg residue at position used as a PTH receptor agonist for the treatment of the above mentioned disorders and fracture repair.
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Pred. No. 3.7e-13;
0; Mismatches 1;
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ID AAR11731 standard; Protein;
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                                                                                                                                               Bringhurst FR, Takasu H,
                                                                                               (GEHO ) GEN HOSPITAL CORP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Conservative
                                                                                                                                                                                               WPI; 2000-400076/34.
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Best Local Similarity
Matches 26; Conserv
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24 - NOV - 1999;
                                               25-NOV-1998;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           cells, initiating signal transduction. It has been identified that the carboxyl terminal of PTH is important for PTH receptor binding, while the amino terminal is important for signal transduction. Various PTH peptides were produces with amino- and carboxy terminal modifications which had warying PTH receptor activation properties and therefore downstream signalling. Aberrant PTH activity has been implicated in a number of disorders: osteoporosis, osteopaenia, hypoparathyroidism and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   and a variety of cancers: breast, lung and prostate carcinoma, multiple myeloma and epidermoid cancers of the head, neck and oesophagus. The myeloma and epidermoid cancers of the head, neck and oesophagus. The present sequence is a PTH peptide, with a Gly residue at position 1 and a Glu residue emilinates downstream signalling via phospholipase C (PLC) and the Glu residue reduces PLC signalling and ligand binding. PTH peptides with an Ala residue at position 1 and an Arg residue at position 19 have improved signalling properties and therefore may be suitable as PTH receptor agonists for the treatment of the above mentioned disorders and fracture repair.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    In turn, hypercalcaemia is associated with hypernephroma
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Parathyroid hormone (PTH) binds to PTH receptors in renal and osseous
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                           Novel biologically active peptide comprising a parathyroid hormone peptide derivative, useful for treating osteoporosis -
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human; parathyroid hormone; signal transduction; osteoporosis; osteopaenia; hyporathyroidism; fracture repair; hypercalcaemia; breast cancer; lung cancer; prostate cancer; multiple myeloma; hypernephroma; head and neck epidermoid cancer; oesophagus cancer;
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0
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                                                                                                                                                                                                                                                                      Potts JT;
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Human parathyroid hormone peptide # 12.

osteopathic; PTH.

WO200031266-A1

02-JUN-2000

Homo sapiens

04-SEP-2000 (first entry)

AAY98052;

28

Sequence

AAY98052 standard; peptide; 28 AA

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Gaps

Gaps

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Indels

Length 29;

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New human parathyroid hormone analogues - which have increased adenylyl cyclase activating activity, used for treating osteoporosis
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAR88829-R88841 are human parathyroid hormone (hPTH) analogues. The
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   analogues increase G-protein coupled adenylyl cyclase (cAMPase) activity and reduce protein kinase C (PKC) activity. The analogues can reverse the loss of bone and increase bone mass and density without undesirable effects. They are useful for the treatment of osteoporosis and other bone related disorders and disorders involving bone cell calcium regulation.
                                                                                                                                                                                                                                                                                                                                                                                                                      Parathyroid hormone; PTH; analogue; osteoporosis; bone cell; calcium regulation; reduced PKC activity; protein kinase C; increased adenylyl cyclase activity; cAMPase; bone loss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 134; DB 17; Length 30;
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                                                                                                                                                                                                                                                                                                                                                                                      Human parathyroid hormone analogue, hPTH(1-30)-NH2.
                                                                      Score 134; DB 17;
Pred. No. 3.8e-13;
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); Mismatches
                                                                                                           0; Mismatches
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30
                                                                                                                                                                    2 vseiglmhnlgkhlnsmervewlrkkl
                                                                                                                                               2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
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0
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                                                                       97.8%;
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                                                                                                                                                                                                                                                                                                                                                  07-OCT-1996 (first entry)
                                                                                                             Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Neugebauer W, Sung WL,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (WILL/) WILLICK G E.
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                                                                                          Best Local Similarity
Matches 26; Conserv
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Best Local Similarity
Matches 26; Conserv
                    AA;
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                  Sequence
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                                                                         Query Match
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                                                                                                                                                                                                                                                                                                                                                    Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAR8829-R88841 are human parathyroid hormone (hPTH) analogues. The analogues increase G-protein coupled adenylyl cyclase (CAMPase) activity and reduce protein kinase C (PKC) activity. The analogues can reverse the loss of bone and increase bone mass and density without undesirable effects. They are useful for the treatment of osseoporosis and other bone related disorders and disorders involving bone cell calcium regulation.
                                                                                                                                             The sequence is encoded by adenine rich codons. Codons 29-84 (See AAQ11618) are degenerate in the usage frequency favoured by E.coli or yeast. A plasmid conty. the complete sequence expresses PTH with an improved yield. PTH is a blood calcium regulator known
                                  Mature human parathyroid synthesis - includes using eg E. coli transformed by plasmid contg. synthetic nucleotide sequence contg. adenine rich codons in N-terminal region.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  New human parathyroid hormone analogues - which have increased
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Parathyroid hormone; PTH; analogue; osteoporosis; bone cell; calcium regulation; reduced PKC activity; protein kinase C; increased adenylyl cyclase activity; cAMPase; bone loss.
                                                                                                                                                                                                                                                                                                                Length 29;
                                                                                                                                                                                                                                                                                                                                                  Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human parathyroid hormone analogue, hPTH(1-29)-NH2
                                                                                                                                                                                                                                                                                                              Score 134; DB 12;
Pred. No. 3.8e-13;
                                                                                                                                                                                                                                                                                                                                                  0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                     3 vseiglmhnlgkhlnsmervewlrkkl 29
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29
                                                                                                                                                                                                                                                                                                                                                                                      2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAR88836 standard; peptide; 29 AA.
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                                                                                                           Disclosure; Fig 3; 62pp; English
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                                                                                                                                                                                                                                                                                                              97.8%;
96.3%;
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Best Local Similarity 96.3
Matches 26; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   /note=
                                                                                                                                                                                                                       to increase bone mass.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (WILL/) WILLICK G E.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WPI; 1996-151754/16.
N-PSDB; AAQ11617
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Key
Modified-site
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                                                                                                                                                                                                                                                           Sequence
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AAR8836
IID AAR8
XX
AC AAR8
XX
DT 07-0
XX
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DE Huma
DE CC AAR8

13

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Gaps

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Indels

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The present sequence represents a human parathyroid hormone (hPTH) (1-31) peptide analogue. The present invention also describes a method for screening peptides for osteogenic activity by subcutaneous injection of a test compound and seeing if a small drop in arterial pressure occurs after a short time. The hPTH peptide analogue can be useful for stimulating bone growth, restoring bone and promoting bone healing, especially treatment of osteoporosis and normal fractures. The hPTH peptide analogue can be administered by injection or inhalation, peptide analogue can alpha-helix in the receptor-binding region of the hormone and increases adenylly cyclase (AC) activity, while
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   cyclisation increases stability against protesses. The screening method, which can be performed in intact female animals, is a quick and simple way of identifying inactive compounds, avoiding the need for long-term, expensive tests on ovariectomised animals.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Cyclic human parathyroid hormone peptide(s) with 27Lys substitution - for treating osteoporosis and fractures, also method for screening osteogenic peptide(s) based on their hypotensive action
                                                                                                                                                                                                                                                                                                               26
/note= "Lys is bound to Glu at position 22 to form
a cyclic structure"
                                                                                                                                                                                                                                                                                   /note= "Glu is bound to Lys at position 26 to form
                                                                                                                                                         Human; parathyroid hormone; hPTH; cyclic; osteoporosis; fracture;
                                                                                                                        Human parathyroid hormone cyclic peptide analogue SEQ ID NO:14.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Whitfield J;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Ross V,
                                                                                                                                                                                                                                                                                                     a cyclic structure"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Morley P, Neugebauer W,
                                                                                                                                                                                                                                                      Location/Qualifiers
                               AAW42059 standard; peptide; 31 AA.
                                                                                                                                                                                                                                                                                                                                                              31
/note= "amidated"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       97US-0040560.
96US-0691647.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          97WO-CA00547.
                                                                                         06-JUL-1998 (first entry)
                                                                                                                                                                           hypotensive action; bone.
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                                                                                                                                                                                                                                                                                                                                                                                                            WO9805683-A1
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02-AUG-1996;
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Willick GE;
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                                                              AAW42059;
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RESULT 15
                 AAW42059
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Gaps

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Length 31; Indels

Score 134; DB 19; Pred. No. 4.1e-13; 0; Mismatches 1;

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Conservative

Query Match Best Local Similarity Matches 26; Conserv

97.8%;

2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28

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2 vseiglmhnlgkhlnsmervewlrkkl 28

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Search completed: August 31, 2001, 16:00:42 Job time: 46 sec

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0991c7 cryctolagus 031521 poecilia re 09km16 vibrio chol 031523 poecilia re 09zu32 arabidopsis P97053 pseudomonas

Q9u8u9 eptatretus Q90269 brachydanio 024430 glycine max Q39016 arabidopsis arabidopsis arabidopsis

Q92U32 P97053 Q9U8U9 Q90269

Q9KNJ6 Q31523

10

Q9GEC7 Q31521

386 758 1048 544 790 1111 121 1124 1177 285 2297 313 323

Q39016 Q38869 Q91qh7

Q42354 arabidopsis Q9gk30 ovis aries Q31524 poecilia re

P79749 fugu rubrip

Q9x3r2 pseudomonas Q9vik6 drosophila 29w0e8 drosophila

P79004 Q9W0E8 P79749 Q9X3R2 Q9VIK6 Q42354 Q9GK30

099cm7 mus musculu 09de49 brachydanio 09puf6 gallus gall 089625 human immun 091177 oryzias lat 002528 oryzias lat

Q9QZM7 Q9DE49

091.QH7

O9PUF6

Q39016 Q38869

02443(

113 113 110 110

401 457 490 495

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MEDLINE=20082971; PubMed=10613847;
Caetano A.R., Shiue Y.L., Lyons L.A., O'Brien S.J., Laughlin T.F.,
Bowling A.T., Murray J.D.;
A comparative and of the horse (Equus caballus).";
Genome Res. 9:1239-1249(1999).
EMBL; AF134233; AAF62347.1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Equus caballus (Horse).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Perissodactyla; Equidae; Equus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               9805 MW; 253184EA681A2022 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                     (TrEMBLrel. 15, Last sequence update)
(TrEMBLrel. 16, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                           (TrEMBLrel. 15, Created)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Pfam; PF01279; Parathyroid; 1.
PROSITE; PS00335; PARATHYROID; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            HORMONE (FRAGMENT)
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Best Local Similarity 92.65
Matches 25; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                   PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               InterPro; IPR001415; -
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 86 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       NCBI_TaxID=9796;
                                                                                                                                                                                                                                                                                                                                                                                                                                                            PARATHYROID
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01-MAR-2001
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SEQUENCE
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Q9GL67
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RESULT
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094167 felis silve
092016 mus musculu
091809 fugu rubrip
091802 sparus aura
091404 drosophila
094046 drosophila
094048 arabidopsis
09149 arropyrum p
09149 arropyrum p
09142 caenorhabdi
090049 lasmodium
025713 helicobacte
09490 caenorhabdi
090009 plasmodium
025713 helicobacte
09490 caenorhabdi
090009 plasmodium
                                                                             August 31, 2001, 15:59:56; Search time 21.89 Seconds (without alignments) 169.234 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Q9pvu7 lampetra re
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Description
                                                                                                                                                                                                                                         425026
           GenCore version 4.5
Copyright (c) 1993 - 2000 Compugen Ltd
                                                                                                                                                       1 XVSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                        Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                 425026 seqs, 132305027 residues
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09GL67
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Maximum Match 100%
Listing first 45 summaries
                                                          protein search, using sw model
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Q9I8E9
Q9I8U2
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024164
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09VLQ8
09UH68
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P91490
Q9U8W5
Q17633
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Gapop 10.0 , Gapext 0.5
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Q9U0H9
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sp_unclassified:*
sp_vertebrate:*
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sp_invertebrate:*
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sp_organelle:*
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sp_bacteria:*
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length: 2000000000
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Match Length DB
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Maximum DB seq
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Length 86; 1; Indels

Score 130; DB 6; Pred. No. 2.5e-12; 1; Mismatches 1.

94.98; 92.68;

115 AA.

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MEDLINE-20304721; PubMed-10843788; Flanagan J.A., Power D.M., Bendell L.A., Guerreiro P.M., Fuentes J., Flanagan J.A., Canario A.V.M., Danks J.A., Brown B.L., Ingleton P.M.; Cloring of the cDNA for sea bream (Sparus aurata) Parathyroid hormone-related protein."; EmBL: Bendorinol. 118:373-382(2000).

EmBL: AF197904; AAF79073.1; -...
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Actinopterygii; Neopterygii; Teleostei; Euteleostei; Neoteleostei;
Acanthomorpha; Acanthopterygii; Percomorpha; Perciformes; Percoidei;
                                                                                                                                                                   Bukaryota; Metazoa; Chordata; Craniata; Verfebrata; Buteleostomi;
Actinopterygii; Neopterygii; Teleostei; Buteleostei; Neoteleostei;
Acanthomorpha; Acanthopterygii; Percomorpha; Tetraodontiformes;
Tetraodontidae; Takifugu.
NCBI_TaxID=31033;
                                                                                                                                                                                                                                                                                           MEDLINE=2011478; PubMed=10854780; Power D.M., Flannigan J., Ingleton P.M., Canario A.V.M., Danks J., Elgar G., Clark M.S.; "Genomic structure and expression of parathyroid hormone-related protein in a teleost, Fugu rubripes."; Gene 250:67-79(2000).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        43.8%; Score 60; DB 13; Length 163; 48.1%; Pred. No. 0.17;
                                                                                                                                                    (Japanese pufferfish) (Takifugu rubripes)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    POTENTIAL.
3AC5F2C764732278 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 -; 1.
: 18722 MW; 6E8D5E07F9E5EDC9 CRC64;
                                                       01-OCT-2000 (TrEMBLrel. 15, Created)
01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
PARATHYROID HORMONE-RELATED PROTEIN PRECURSOR.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-0CT-2001 (TrEMBLrel. 16, Last annotation update)
PARAPHYROLD HORMONE-RELATED PROFELIN PRECURSOR.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              162 AA
                      163 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            sea bream)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        34 Po
18698 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           InterPro; IPR003571; -.
Pfam; PF01279; Parathyroid; 1.
                                                                                                                                                                                                                                                                                                                                                                                                           CAB94712.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                Pfam; PF01279; Parathyroid; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                13; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                               IPR001415;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SIĞNAL 1
SEQUENCE 163 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                SEQUENCE FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                             EMBL; AJ249391;
                                                                                                                                                    Fugu rubripes
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                                                                                                                                                                                                                                                                                                                                                                                                                                InterPro;
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                                       Q918E9;
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STRAIN-129/SV;
Karaplis A.C., He B., Hiou-Tim F.F.T., Al-Akad B., Kronenberg H.M.;
"Cloning of the murine gene encoding parathyroid hormone: genomic organization and nucleotide sequence.";
Submitted (MAY-1998) to the EMBL/GenBank/DDBJ databases.
EMBL, AF066075; AAC99656.1;
HSSP: PO1270: 12WB.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Mus musculus (Mouse).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Musinae; Mus
NCBI_TaxID=10090;
                                                                                               Euteleostomi;
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TOTIBLE R.E., Kohn C.W., Leone G.W., Capen C.C., Rosol T.J.;
"Molecular cloning of feline preproparathyroid hormone.";
Submitted (OCT-2000) to the EMBL/GenBank/DDBJ databases.
EMBL; AF309967; AAG30545.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 115;
                                                                                                                                                                                                                                                                                                                                                                     ch 86.1%; Score 118; DB 6; Length 115; 1 Similarity 81.5%; Pred. No. 2.2e-10; 22; Conservative 3; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Indels
                                                                                                                                                                                                                                                                              POTENTIAL.
PARATHYROID HORMONE.
; 80CD557CC6A1A47E CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PARATHYROID HORMONE.
DA43FABBCB4E2FD9 CRC64;
                                                                      Felis silvestris catus (Cat).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata;
Mammalia; Eutheria; Carnivora; Fissipedia; Felidae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (TrEMBLrel. 10, Created)
(TrEMBLrel. 10, Last sequence update)
(TrEMBLrel. 16, Last annotation update)
01-MAR-2001 (TrEMBLrel. 16, Last sequence update) 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       85.4%; Score 117; DB 11;
81.5%; Pred. No. 3.1e-10;
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                                      PREPROPARATHYROID HORMONE PRECURSOR
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ProDom; PD010687; -; 1.
PROSITE; PS00335; PARATHYROID; 1.
                                                                                                                                                                                                                                                                              25 PO
115 PA
12921 MW;
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115 PA
12825 MW;
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CHAIN 32 1
SEQUENCE 115 AA;
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                                                                                                                               NCBI_TaxID=9685;
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CHAIN 32
SEQUENCE 115 7
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Query Match

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Score 56; DB 13; Length 162; Pred. No. 0.67; 40.98; Best Local Similarity

Query Match

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Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha; Ephydroidea; Drosophilidae; Drosophila.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Pterygota: Neoptera: Endopterygota: Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
NCBI_TaxID=7227;
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"Toll-related receptors and the control of antimicrobial peptide expression in Drosophila.";
Proc. Natl. Acad. Sci. U.S.A. 0:0-0(2000).
EMBL; AF247769; AAF86229.1;
Flybase: FBD00015770; MstProx.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Drosophila melanogaster (Fruit fly).
Eukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            83175 MW; 007CD7AEF67BC1B0 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                        01-NOV-1996 (TrEMBLrel. 01, Last sequence update) 01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
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01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    785 AA
                                                                                                          213 INEEQLLQSMHRKLNNLNTIMSIYKYMEWLHRKL 246
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  36.9%; Score 50.5; D
29.4%; Pred. No. 22;
Live 10; Mismatches
Mismatches
                                                               2 VSEIQLMHNLGKHLNSMXRV-----EWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        2 VSEIQLMHNLGKHLNSMXRV-----EWLRKKL 28
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              interleukin-1 receptor family.";
J. Biol. Chem. 271:5777-5783(1996).
EMBL; U42425; AAC46999.1; -.
10;
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   Conservative
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                                                                                                                                                                                                                                                                                           PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            InterPro; IPR001611; -. Pfam; PF00560; LRR; 2. Pfam; PF01462; LRRNT; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Pfam; PF01582; TIR; ]
SMART; SM00255; TIR;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                InterPro; IPR000157;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             InterPro; IPR000157;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
Matches 10; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            711 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     MSTPROX OR CG1149.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     MSTPROX OR CG1149
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SEQUENCE FROM N.A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TOLL-3 (FRAGMENT)
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10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                              DMMSTPROX.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SEQUENCE
                                                                                                                                                                                                                                                                                                                         024164;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       09NBK6
                                                                                                                                                                                                                                                                                           024164
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      Matches
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Q24164
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RAM Addams M.D., Celniker S.E., Holf R.A., Evans C.A., Gocayne J.D.,
RA Adams M.D., Celniker S.E., Holf R.A., Evans C.A., Gocayne J.D.,
RA Amanatides P.G., Scherer S.E., Li P.W., Hoskins R.A., Galle R.F.,
RA Gacrage R.A., Lewis S.E., Richards S., Ashburner M., Henderson S.N.,
RA Sutton G.G., Wortman J.R., Yandell M.D., Zhang Q., Chen L.X.,
RA Brandon R.C., Ragers Y. H.C., Blazej R.G., Champe M., Pfeiffer B.D.,
RA Man K.H., Doyle C., Baxter B.G., Helt G., Nelson C.R., Miklos G.L.G.,
RA Abril J.F., Agbayani A., An H.-J., Andrews-Pfannkoch C., Baldwin D.,
RA Ballew R.M., Basu A., Baxendale J., Bayraktaroglu L., Beasley E.M.,
RA Berson K.Y., Benos P.V., Berman B.P., Bhandari D., Bolshakov S.,
RA Burtis K.C., Busam D.A., Bulter H., Cadieu E., Center A., Chandra I.,
RA Burtis K.C., Busam D.A., Bulter H., Cadieu E., Center A., Chandra I.,
RA Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
RA Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
RA Cherry J.M., Cawley S., Dahlke C., Bowenport L.B., Davies P.,
RA Charry J.M., Cawley S., Carlier A., Changel S., Dunkov B.C., Dunn P.,
RA Horlin K.J., Evangelista C.C., Ferraz C., Ferriara S., Fleischmann M.,
R. Fosler C., Gabriellan A.E., Garg N.S., Gelbart W.M., Glasser K.,
RA Hostin D., Houston K.A., Harnandez J.R., Houck J.,
RA Hostin D., Houston K.A., Hayland T.J., Well M.-H., Ibegwan C.,
RA Kimmel B.E., Kodira C.D., Kraft C., Kravitz S., Kulp D., Lai Z.,
RA Liu X., Mattei B., McIntosh T.C., McLeod M.P., McPherson D.,
RA Melson D.R., Nelson K.A., Nixon K., Nixon B.,
Ralazolo M., Pittman G.S., Pan S., Pollard J., Puri V., Reese M.G.,
RA Ralazolo M., Pittman G.S., Pan S., Pollard J., Puri V., Ranith T.,
RA Spier E., Spradling A.C., Stapleton M., Strong R., Santh T.,
RA Man Z.-Y., Wangarman D.A., Wenter E., Wang A.H.,
RA Wang Z.-Y., Wangarman D.A., Wenter E., Wang A.H.,
RA Wang Z.-Y., Wang A., There R., Venter E., Wang A., Wang Z.-Y., Wang A., There R., There R., There R., Wang Z.-Y. Wang Z.-P., Siden K., Pacier E., Siden K., Pacier E., Salen H.,
RA W
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"The genome sequence of Drosophila melanogaster.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
Ephydroidea; Drosophilidae; Drosophila.
      Gaps
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      Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 57BA402F6AF764D5 CRC64;
                                                                                                                                                                                                                                                                                                                                             01-MAY-2000 (TrEMBLrel. 13, Created)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
MSTPROX PROTEIN.
      12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 50.5; DB 5;
Pred. No. 8.8;
                                                                                                                                                                                                                                                                                              303 AA
      Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MSTPROX OR CG1149.
Drosophila melanogaster (Fruit fly).
                                                                     2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                          MEDLINE=20196006; PubMed=10731132;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               36.9%;
      Conservative
                                                                                                                                                                                                                                                                                                 PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Pfam; PF01462; LRRNT; 1
SMART; SM00013; LRRNT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       InterPro; IPR000372
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Quary Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SEQUENCE FROM N.A.
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      12;
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Gaps

7;

Length 711; Indels

5;

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Query Match
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Q9LH68
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Bukaryota; Metazoa; Arthropoda; Tracheata; Hexapoda; Insecta;

Pterygota; Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;

Ephydroidea; Drosophilidae; Drosophila.
                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MEDLINE-20196006; PubMed-10731132; Adams M.D., Celniker S.E., Holt R.A., Evans C.A., Gocayne J.D., Amanatides P.G., Scherer S.E., Li P.W., Hoskins R.A., Galle R.F., George R.A., Lewis S.E., Richards S., Ashburner M., Henderson S.N., Sutton G.G., Wortman J.R., Yandell M.D., Zhang Q., Chen L.X., Brandon R.C., Rogers Y.-H.C., Blazej R.G., Champe M., Pfelffer B.D.,
                                                                                                                                                                                                                                                                                                                                                                       7;
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                                                                                                                                                                                                                                                                                                                 Length 785;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 36.5%; Score 50; DB 2; Length 336, 42.3%; Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                       Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11; Indels
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STRAIN-C-125 / JCM 9153;
Takami H., Nakasone K., Takaki Y.;
Submitted (MAR-2000) to the EMBL/GenBank/DDBJ databases.
EMBL; AP001518; BAB06973.1; -
SEQUENCE 336 AA; 34615 MW; B661697C95EA6736 CRC64;
                                                                                                                                                                                                              785 AA; 91897 MW; C8EFA786D7CA9525 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Bacteria; Firmicutes; Bacillus/Clostridium group;
Bacillus/Staphylococcus group; Bacillus.
                                                                                                                                                                                                                                                                                                              5;
                                                                                                                                                                                                                                                                                                              DB
                                                                                                                                                                                                                                                                                                       ; Score 50.5; DE; Pred. No. 24; 10; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     336 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ::| ||: :: :||:: :|| :1| 287 INEEQLLQSMHRKLNNLNTIMSIYKYMEWLHRKL 320
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              4; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                2 VSEIQLMHNLGKHLNSMXRV-----EWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Created)
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                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity 29.4%;
Matches 10; Conservative 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        01-OCT-2000 (TrEMBLrel. 15, 01-OCT-2000 (TrEMBLrel. 15, 01-OCT-2000 (TrEMBLrel. 15,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               01-MAY-2000 (TrEMBLrel. 13, 01-MAY-2000 (TrEMBLrel. 13, 01-MAR-2001 (TrEMBLrel. 16,
                                                  Pfam; PF00560; LRR; 2.
Pfam; PF01462; LRRNT; 1.
Pfam; PF01582; TIR; 1.
SMART; SM00013; LRRNT; 1.
NON_TER
SEQUENCE 785 AA; 91897 M
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nes 11; Conservative
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   InterPro;
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                                  InterPro
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Q9K7V5;
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10 09877V5

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RESULT

RESULT
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1009VLQ8
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RA Abril J.F., Agbayani A., An H.-J., Andrews-Pfannkoch C., Baldwin D., Ballew R.M., Basu A., An H.-J., Andrews-Pfannkoch C., Baldwin D., Ballew R.M., Basu A., An H.-J., Andrews-Pfannkoch C., Baldwin D., Ballew R.M., Basu A., An H.-J., Andrews-Pfannkoch C., Baldwin D., Ballew R.M., Basu A., Baxendale J., Bayraktaroglu L., Beasley E.M., Becson K.Y., Bencos P.V., Bernan B.P., Bhandari D., Bolshakov S., Bortchan M.R., Bouck J., Brokstein P., Brottler P., Butter H., Cawley B.M., Chandra I., Burtis K.C., Busam D.A., Daller H., Davenport L.B., Davies P., Danner D., Delcher A., Deng Z., Mays A.D., Dew I., Dieter S.M., Dadson K., Doup L.E., Downes M., Dugan-Rocha S., Dunkov B.C., Dun D. B., Downes M., Dugan-Rocha S., Dunkov B.C., Dun D. B., Downes M., Dugan-Rocha S., Dunkov B.C., Dun D. B., Andrei B. C., Gabrielian A.E., Garg N.S., Gelbart W.M., Glasser K., Glodek A., Gong F., Correll J.H., Gu Z., Guan P., Harris M., Harris M., Haush F., Karpen G.H., Ke Z., Kennison J.A., Ketchum K.A., Haush F., Karpen G.H., Ke Z., Kennison J.A., Ketchum K.A., Jalali M., Kalush F., Karpen G.H., Ke Z., Kennison J.A., Ketchum K.A., Jasko P., Lei Y., Levitsky A.A., Li J., Liz Z., Liang Y., Lin X., Liu X., Mattei B., McIntosh T.C., McLeod M.P., McPherson D., Mount S.M., Mullshinan N.V., Mobarry C., Moriss J., Mosherson D., Mount S.M., Mullshinan N.V., Nobsarry C., Moriss J., Mosher B., Spradling A.C., Stapleton M., Stupski M.P., Shul H., Shult H., Sprad Spradling A.C., Stapleton M., Stupski M.P., Smith T., Williams S.M., Woodage T., Worley K.C., Wu D., Yang S., Yao Q., Zheng L., Wu D., Yang S., Yao Q., Zheng L., The Genome Sequence of Drosophila melanogaster.", The Genome Sequence of Drosophila melanogaster.", Education R. Schole R. A., Mychar B., Ruby M., Woodage T., Worley K.C., Wu D., Yang S., Yao Q., Zheng L., Shong R., The Genome Sequence of Drosophila melanogaster.", Education R. S., Mang S., Zhu X., Zhu X.,
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Eukaryota; Viridiplantae; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae; eurosids II; Brassicaeses; Arabidopsis.
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ATP-binding; Transferase; Tyrosine-protein kinase.
SEQUENCE 1500 AA; 169216 MW; 4B4FBD5DAA505B5E CRC64;
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Last annotation update)
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60.0%; Pred. No. 56;
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PROSITE; PS00107; PROTEIN_KINASE_ATP; 1.
PROSITE; PS50011; PROTEIN_KINASE_DOM; 1.
PROSITE; PS00109; PROTEIN_KINASE_IYR; 1.
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FlyBase; FBgn0032006; CG8222.
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Pfam; PF00069; pkinase; 2.
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Matches 9; Conservative
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907 VSELKIMVHLGQHLN 921
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SEQUENCE FROM N.A
                                                                STRAIN-BRISTOL N2
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Kawarabayasi Y., Hino Y., Horikawa H., Yamazaki S., Haikawa Y.,
Jin-no K., Takahashi M., Sekine M., Baba S.-I., Ankai A., Kosugi H.,
Jin-no K., Takahashi M., Sekine M., Makazawa H.,
Takamiya M., Masuda S., Funahashi T., Tanaka T., Kudoh Y.,
Yamazaki J., Xushida N., Oguchi A., Aoki K. II., Kubota K.,
Makamura Y., Nomura N., Sako Y., Kikuchi H.,
"Complete genome sequence of an aerobic hyper-thermophilic
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    Rhabditidae; Peloderinae; Caenorhabditis.
    NCBL_TaxID=6239;
                                                                                                                          Nakamura Y.;
"Structural Analysis of Arabidopsis thaliana Chromosome 3. III.";
"Structural Analysis of Arabidopsis thaliana Chromosome 3. III.";
Submitted (MAY-2000) to the EMBL/GenBank/DDBJ databases.
EMBL; AP002062; BAB02553.1; -.
SEQUENCE 63 AA; 7025 MW; 2C8966DDEB4673DD CRC64;
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Archaea; Crenarchaeota; Desulfurococcales; Desulfurococcaceae;
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                                                                                                                                                                                                                                                                                                                                                          Score 49; DB 10; Length 63; Pred. No. 2.8; 4; Mismatches 5; Indels
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SEQUENCE 115 AA; 12603 MW; D9A02D3E1D5CC232 CRC64;
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01-NOV-1999 (TrEMBLrel. 12, Last sequence update)
01-MAR-2001 (TrEMBLrel. 16, Last annotation update)
HYPOTHETICAL 12.6 KDA PROTEIN APE1063.
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Last annotation update)
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DNA Res. 6:83-101(1999).
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47.1%;
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01-0CT-2000 (TrEMBLrel. 15,
01-MAR-2001 (TrEMBLrel. 16,
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Best Local Similarity 47.1:
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                                                                SEQUENCE FROM N.A.
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                                                                                                      STRAIN=COLUMBIA;
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                                       "Genome sequence of the nematode C. elegans: a platform for investigating biology. The C. elegans Sequencing Consortium."; Science 282:2012-2018(1998).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Eukaryota; Alveolata; Apicomplexa; Haemosporida; Plasmodium.
NCBL_TaxID=36329;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 244;
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Devlin K., Bowman S., Churcher C., Harris B., Harris D., Quail M., Barrell B.;
Submitted (FBE-1999) to the EMBL/GenBank/DDBJ databases.
EMBL, AL035475; CAB62884.1; ...
Hypothetical protein.
SEQUENCE 977 AA; 117527 MW; 3FBE5600D7A0F61D CRC64;
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                                                                                                                                                                                                                                                                                                                                  Submitted (WAR-2000) to the EMBL/GenBank/DDBJ databases. EMBL; AC006795; AAF59493.1; -. InterPro; IPR001719; -. InterPro; IPR001245; -. Pfam: PF00169; pkinase; 1. PRNIF; PR00109; TYRKINASE. PROSTITE; PS0011; PROTEIN KINASE. DOM; 1. PROSTITE; PS00109; PROTEIN_KINASE_TYR; UNKNOWN_1.
                                                                                                                                                                                                  "The sequence of C. elegans cosmid Y50D4B.";
Submitted (MAR-1999) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       SMART; SM00219; Tyrkc; 1.
ATP-binding; Transferase; Tyrosine-protein kinase.
SEQUENCE 244 AA; 28568 MW; C389C56F7E31524A CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  01-WAY-2000 (TrEMBLrel. 13, Created)
01-WAY-2000 (TrEMBLrel. 13, Last sequence update)
01-WAR-2001 (TrEMBLrel. 16, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 5;
51;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ore 49; DB 5
ed. No. 12;
Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          977 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       35.8%; Score 49; DB
33.3%; Pred. No. 51;
tive 9; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 49;
Pred. No. 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Plasmodium falciparum (isolate 3D7).
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MEDLINE=99069613; PubMed=9851916;
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Best Local Similarity 33.3
Matches 8; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match 35.8
Best Local Similarity 36.4
Matches 8; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PRELIMINARY;
                                                                                                                                   SEQUENCE FROM N.A.
STRAIN-BRISTOL N2;
Lamar B., Le T.;
                                                                                                                                                                                                                                                                      SEQUENCE FROM N.A. STRAIN=BRISTOL N2;
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025713

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ID 025713 PRELIMINARY; PRT; 207 AA.

C 025713; PRELIMINARY; PRT; 207 AA.

DT 01-JAN-1998 (TrEMBLE-1. 05, Last sequence update)

DT 01-JAN-1998 (TrEMBLE-1. 14, Last annotation update)

DE HYPOTHETICAL 23.7 KDA PROTEIN.

SHIGH HIGH.

HIGH HIGH.

C Bacteria; Proteobacteria; epsilon subdivision; Helicobacter group;

C Bacteria: Proteobacteria; epsilon subdivision; Helicobacter group;

RA Potton M.D.; Weldardson D.; Dodoson R., Klandak H.G., Glodek A.,

RA McKenney K., Fitzgerald L.M., Lee N., Adams M.D., Hickey J.M.,

RA McKenney K., Fitzgerald L.M., Karp P.D., Smith H.O., Fraser C.M.,

RA Potton M.D., Weldman J.M., Fujii C., Bowman C., Watthey L., Wallin E.,

RA Venter J.C.;

RY The complete genome sequence of the gastric pathogen Helicobacter

RY The Complete genome sequence of the gastric pathogen Helicobacter

RY The RABUS, RABUS, AAD08133.1; -.

DR REMEL, RABUSOR13.1; -.

KW Hypothetical protein.
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Search completed: August 31, 2001, 16:01:30 Job time: 94 sec

Gaps

Score 47; DB 2; Length 207; Pred. No. 20; 2; Mismatches 7; Indels

Ouery Match 34.3%; Best Local Similarity 50.0%; Matches 9; Conservative

οy

us-09-448-867-1.rsp

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GenCore version 4.5
Copyright (c) 1993 - 2000 Compugen Ltd.
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OM protein - protein search, using sw model

August 31, 2001, 15:59:56; Search time 10.02 Seconds (without alignments) 95.724 Million cell updates/sec Run on:

US-09-448-867-1 137 1 XVSEIQLMHNLGKHINSMXRVEWLRKKL 28

Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

93435 seqs, 34255486 residues Searched:

93435 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

SwissProt_39:* Database :

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

	Description	P01270 homo sapien	Q9xt35 macaca fasc	P01269 sus scrofa	P01268 bos taurus	P52212 canis famil	P04089 rattus norv	-	thermo	051708 borrelia bu		-		P05622 mus musculu	P09619 homo sapien	-	P17251 gallus gall		P12272 homo sapien	P13085 rattus norv	P02793 rattus norv	P39917 coxiella bu	Q09189 schizosacch			P26619 xenopus lae		P16234 homo sapien	P26618 mus musculu	_	_	Q01742 homo sapien	uns mus	P18460 gallus gall
SUMMARIES	Ð	PTHY_HUMAN	PTH_MACEA	PTHY_PIG	PTHY_BOVIN	PTHY_CANFA	PTHY_RAT	PTHY_CHICK	SYL_THEMA	MURG_BORBU	GAF2_SCHPO	CDPK_SOYBN	TORA_ECOLI	PGDR_MOUSE	PGDR_HUMAN	PTHR_MOUSE	PTHR_CHICK	PTHR_CANFA	PTHR_HUMAN	PTHR_RAT	FRIL_RAT	LOLA_COXBU	AZR1_SCHPO	PYRG_HELPJ	PYRG_HELPY	PGDS_XENLA	PGDS_RAT	PGDS_HUMAN	PGDS_MOUSE	YJCC_ECOLI	MERA_BACSR	BFR2_HUMAN	FGR3_MOUSE	CEK2_CHICK
	° Query Match Length DB	115 1			115 1				824 1	363 1	564 1	508 1	848 1	1098 1	1106 1	175 1	176 1	177 1	177 1	177 1	182 1	211 1	224 1	538 1	538 1	1087 1	1088 1	1089 1	1089 1	528 1	631 1	654 1	801 1	806 1
ð	Query Match	97.8	97.8	92.0	91.2	91.2	84.7	65.7	39.4	38.0	37.2	36.9	36.5	34.3	34.3	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.6	33.2	32.8	32.8	32.8	32.8
	Score	134	134	126	125	125	116	06	54	52	51	50.5	20	47	47	46	46	46	46	46	46	46	46	46	46	46	46	46	46	45.5	45	45	45	45
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P22607 homo sapien Q03364 xenopus lae	P21802 homo sapien P21803 mus musculu	P18461 gallus gall P37285 rattus norv	Q07866 homo sapien P29391 mus musculu	P49945 mus musculu O9xd74 rhizobium m	P09350 physarum po P74309 synechocyst	
FGR3_HUMAN FGR2_XENLA	FGR2_HUMAN FGR2_MOUSE	CEK3_CHICK	KNLC_HUMAN FRL1 MOUSE	FRL2_MOUSE	SR1A_PHYPO ALF1_SYNY3	
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806 813	821 821	823	569	182	300	
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45 45	45	45	44.5	4 4	77	
34 35	36 37	38	40	4 4 3	44	

ALIGNMENTS

RESULT 1 TRESULT 1 TO THE STATE 1 AC PO127 AC PO127 BD THY HUMAN 1 13AUD DT 13

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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      the Swiss Institute of Bioinformatics and the EMBL outstation
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J. Clin. Invest. 86:1084-1087(1990).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- DISEASE: DEFECTS IN PTH ARE A CAUSE OF FAMILIAL ISOLATED
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Klaus W., Dieckmann T., Wray V., Schomburg D., Wingender E., Mayer "Investigation of the solution structure of the human parathyroid hormone fragment (1·34) by 1H NMR spectroscopy, distance geometry, and molecular dynamics calculations."; Biochemistry 30:6936-6942(1991).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Marx U.C., Adermann K., Bayer P., Forssmann W.-G., Rosch P., "Solution structures of human parathyroid hormone fragments hyprH(1-34) and hPTH(1-34) and bovine parathyroid hormone fragment home/1-27, ".
                                                                                                                                                                                                                          Tregear G.W., van Rietschoten J., Green E., Niall H.D., Keutmann H.T., Parsons J.A., O'Riordan J.L.H., Potts J.T. Jr.; "Solid-phase synthesis of the biologically active N-terminal 1-34
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Marx U.C., Austermann S., Bayer P., Adermann K., Ejchart A., Sticht H., Walter S., Schmid F.-X., Jaenicke R., Forssmann W.-G.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MEDLINE-93345518; PubMed-8344299;
Barden J.A., Cuthbertson R.M.;
"Stabilized NMR structure of human parathyroid hormone(1-34).";
Eur. J. Blochem. 215:315-321(1993).
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J. Biol. Chem. 270:15194-15202(1995).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MEDLINE-91009811; PubMed-2212001;
Arnold A., Horst S.A., Gardella T.J., Baba H., Levine M.A..
Kronenberg H.M.;
                                                                                 Keutmann H.T., Niall H.D., O'Riordan J.L.H., Potts J.T. Jr "A reinvestigation of the amino-terminal sequence of human
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         -! - SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              'Mutation of the signal peptide-encoding region of the
                                                                                                                                                                                                                                                                                         peptide of human parathyroid hormone.";
Hoppe-Seyler's Z. Physiol. Chem. 355:415-421(1974).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Biochem. Biophys. Res. Commun. 267:213-220(2000)
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MEDLINE=20090619; PubMed=10623601;
                                                                                                                                                                                                     MEDLINE=75059220; PubMed=4474131;
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Helv. Chim. Acta 56:470-473(1973)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MEDLINE=91299748; Pubmed=2069952;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      MEDLINE=95318084; PubMed=7797503;
                                                              MEDLINE=75146516; PubMed=1125201;
                                                                                                                                              Biochemistry 14:1842-1847(1975).
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                                         REVISIONS
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Malaivijitnond S:, Takenaka 0.; "Nucleotide sequences of parathyroid gene in five species of macaque of Thailand.";
                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            J. Sci. Res. Chulalongkorn Univ. 23:135-142(1998).
--- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN BONE AND PREVENTING THEIR RENAL EXCRETION.
---- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                                                                                                                                                                                                  PARATHYROID HORMONE.
C -> R (IN FIH; LEADS TO INEFFICIENT
PROCESSING OF THE PRECURSOR).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
                                                                                                                                                                                                                                                                                                                                                                                                                          0;
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Pred. No. 2.3e-13;
); Mismatches 1; Indels
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N -> D (IN REF. 5).
849015736A6E5597 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Pfam; PF01279; Parathyroid; 1.
PROSITE; PS00335; PARATHYROID; 1.
Hormone; Signal; Disease mutation; 3D-structure.
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send an email to license@isb-sib.ch).
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          EMBL; J00301; AAA60215.1; -.
EMBL; V00597; CAA23843.1; -.
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NCBI_TaxID=9541;
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                                                                                1HPH; 10-JUL-95.
1LWH; 15-OCT-97.
1ZWB; 12-MAR-97.
1ZWB; 12-MAR-97.
1ZWB; 12-MAR-97.
1ZWF; 16-JUN-97.
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14-JAN-00.
14-JAN-00.
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115 #
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Best Local Simil
Matches 26; (
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Q9XT35;
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                                          Hormone;
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Biochemistry 13:1994-1999(1974).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN
BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MEDIJUE=76018954; PubMed=1164500;
Chu L.L.H., Huang W.-Y., Littledike E.T., Hamilton J.W., Cohn D.V.;
"Porcine proparathyroid hormone. Identification, biosynthesis, and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE OF 32-115.
MEDLINE=74253317; PubMed=4840833;
Sauer R.T., Niall H.D., Hogan M.L., Keutmann H.T., O'Riordan J.L.H.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Eukaryota, Metizoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Cetartiodactyla, Suina, Suidae, Sus.
NCBI_TaxID=9823;
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                                                                                                                                                                                                                                                                 Length 115;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Schmelzer H.-J., Gross G., Widera G., Mayer H.;
Nucleotide sequence of a full-length cDNA clone encoding
Preproparathyroid hormone from pig and rat.";
Nucleic Acids Res. 15:6740-6740(1987).
                                                                                                                                                                                                                                                                                                       Indels
                                                                                                                                                                                                   8C2500EF24BE5597 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           21-UUL-1986 (Rel. 01, Created)
01-UAN-1988 (Rel. 06, Last sequence update)
15-UUL-1998 (Rel. 36, Last annotation update)
PARATHYROID HORMONE PRECURSOR (PARATHYRIN) (PTH).
                                                                                                                                                                                   PARATHYROID HORMONE.
                                                                                                                                                                                                                                                            97.8%; Score 134; DB 1;
96.3%; Pred. No. 2.3e-13;
Live 0; Mismatches 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            115 AA
                                                                                                                                           BY SIMILARITY.
BY SIMILARITY.
                                                                                                                                                                                                                                                                                                                                               VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                    33 VSEIQLMHNLGKHLNSMERVEWLRKKL 59
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SEQUENCE FROM N.A. MEDLINE-87316938; PubMed=3628009;
                                                      InterPro; IPR001415; -. Pfam; PF01279; Parathyroid; 1. PROSITE; PS00335; PARATHYROID; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               partial amino acid sequence.";
Biochemistry 14:3631-3635(1975).
                                                                                                                                      25 B3
31 B3
115 P2
12890 MW;
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                EMBL; AF130257; AAD42777.1;
                                                                                                                                                                                                                                                                                                         Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            STANDARD;
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InterPro; IPR001415;
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PIR; B26806; B26806.
                                                                                                                                                                                                   115 AA;
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                                                                                                                                                                                                                                                                                 Local Similarity
nes 26; Conserv
                                        HSSP; P01270; 1HPY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     scrofa (Pig).
                                                                                                                      Hormone; Signal
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SEQUENCE OF 26-115.
MEDLINE-74142666; PubMed-4522780;
Hamilton J.W., Niall H.D., Jacobs J.W., Keutmann H.T., Potts J.T. Jr.,
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MEDILINE-80056617; PubMed-388425;
Kronenberg H.M., McDevitt B.E., Majzoub J.A., Nathans J., Sharp P.A.,
Ports J.T. Jr., Rich A.,
"Cloning and nucleotide sequence of DNA coding for bovine
                                                                                                                                                                                                                                                                                                             Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MEDIINE-83105964; PubMed=6185374;
Weaver C.A., Gordon D.F., Kemper B.,
"Nucleotide sequence of bovine parathyroid hormone messenger RNA.";
Mol. Cell. Endocrinol. 28:411-424(1982).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea; Bovinae; Bos. NCBI_TaxID=9913;
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Weaver C.A., Gordon D.F., Kissil M.S., Mead D.A., Kemper B.;
"Isolation and complete nucleotide sequence of the gene for bovine
                                                                                                                                                                                                                                                                                                             ö
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MEDLINE-71076162; PubMed-5531031;
Niall H.D., Keutmann H.T., Sauer R., Hogan M.L., Dawson B.F.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  at the 5' terminus of the sense strand of bovine parathyroid
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                                                                                                                                                                                                                                                        Length 115;
                                                                               25
31
115 PARATHYROID HORMONE.
12852 MW; 9FEBBCDE614BAC16 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
30-MRY-2000 (Rel. 39, Last annotation update)
PARATHYROID HORMONE PRECURSOR (PARATHYRIN) (PTH).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 preproparathyroid hormone.";
Proc. Natl. Acad. Sci. U.S.A. 76:4981-4985(1979)
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                                                                                                                                                                                                                                             Score 126; DB 1;
Pred. No. 3.7e-12;
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                                                                                                                                                                                                                                                                                                             2; Mismatches
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Pfam; PF01279; Parathyroid; 1.
PROSITE; PS00335; PARATHYROID; 1.
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MEDLINE=82037785; PubMed=6170060;
                                                                                                                                                                                                                                                  92.0%;
88.9%;
                                                                                                                                                                                                                                                                                                             24; Conservative
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                                                                                                                                      32 1
115 AA;
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                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
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Canis familiaris (Dog).
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                                                                                                                                           MEDLINE=71091588; PubMed-4322265; Potts J.T. Jr., Tregear G.W., Keutmann H.T., Niall H.D., Sauer R., Deftos L.J., Dawson B.F., Hogan M.L., Aurbach G.D.; "Synthesis of a biologically active N-terminal tetratriacontapeptide proc. Natl. Acad. Sci. U.S.A. 68:63-67(1971).
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                                                                                                                                                                                                                                                                                                   Biochem. Biophys. Res. Commun. 267:213-220(2000).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                                                                                                                                                Marx U.C., Aderman R., Bayer P., Forssmann W.-G., Rosch P.; "Solution structures of human parathyroid hormone fragments hPTH(1-34) and hPTH(1-39) and bovine parathyroid hormone fragment bPTH(1-37).";
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            Aurbach G.D., Potts J.T. Jr.;
"The amino acid sequence of bovine parathyroid hormone I.";
Hoppe-Seyler's Z. Physiol. Chem. 351:1586-1588(1970).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            91.2%; Score 125; DB 1; Length 115; 88.9%; Pred. No. 5.2e-12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2; Indels
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V -> G (IN REF. 4).
2ED246B348880710 CRC64;
                                                                      MEDLINE=71063634; PubMed=5275384; Brewer H.B. Jr., Ronan R.; Bovine parathyroid hormone: amino acid sequence."; Proc. Natl. Acad. Sci. U.S.A. 67:1862-1869(1970).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 01-OCT-1996 (Rel. 34, Created)
01-OCT-1996 (Rel. 34, Last sequence update)
15-JUL-1998 (Rel. 36, Last annotation update)
PARATHYROID HORMONE PRECURSOR (PARATHYRIN) (PTH).
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                                                                                                                                                                                                                              STRUCTURE BY NMR OF 32-68.
MEDLINE=20090619; PubMed=10623601;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Pfam; PF01279; Parathyroid; 1.
PROSITE; PS00335; PARATHYROID; 1.
Hormone; Signal; 3D-structure.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        115 AA; 12980 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         EMBL; V00106; CAA23439.1; -. EMBL; J00024; AAA30747.1; -. EMBL; K01938; AAA30749.1; -. EMBL; M25082; AAA30748.1; -.
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InterPro; IPR001415;
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PIR; A24949; A24949.
                                                                                                                                   SYNTHESIS OF 32-65.
                                                           SEQUENCE OF 32-115.
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PTHY_CANFA
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Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
                                                                                                                                                         TISSUE-Parathyroid;
MEDLINE=95369696; PubMed=7642102;
ROSOI T.J., Steinmeyer C.L., McCauley L.K., Groene A.,
BOWILLS J.M., Capen C.C.;
"Sequences of the cDNAs encoding canine parathyroid hormone-related protein and parathyroid hormone.";
Gene 160:241-243(1995).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
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MEDLINE=84135846; PubMed=6321505;
Heinrich G., Kronenberg H.M., Potts J.T. Jr., Habener J.F.;
"Gene encoding parathyroid hormone. Nucleotide sequence of the rat gene and deduced amino acid sequence of rat preproparathyroid
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis.
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MEDLINE=87316938; PubMed=3628009;
MEDLINE=H.-J., Gross G., Widera G., Mayer H.;
Schmelzer H.-J., Gross G., Fill-length cDNA clone encoding preproparathyroid hormone from pig and rat.";
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01-NOV-1986 (Rel. 03, Created)
01-NOV-1986 (Rel. 03, Last sequence update)
01-NOV-1900 (Rel. 40, Last annotation update)
PARATHYROID HORMONE PRECURSOR (PARATHYRIN) (PTH).
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88.9%; Pred. No. 5.2e-12;
tive 1; Mismatches 2;
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BY SIMILARITY.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     33 VSEIQFMHNLGKHLSSMERVEWLRKKL 59
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      InterPro; IPR001415; -Pfam; PF01279; Parathyroid; 1. PROSITE; PS00335; PARATHYROID; 1. Hormone; Signal.
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Best Local Similarity
Matches 24; Conserv
                                                                                                                                 SEQUENCE FROM N.A
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                                                                NCBI_TaxID=9615;
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STRAIN=WSB8 / DSM 3109;
STRAIN=WSB8 / DSM 3109;
MFDLINE-99287316; PubMed-10360571;
Nelson K.E., Clayton R.A., Gill S.R., Gwinn M.L., Dodson R.J.,
Haft D.H., Hickey E.K., Peterson J.D., Nelson W.C., Ketchum K.A.,
McDonald L., Utterback T.R., Malek J.A., Linher K.D., Garrett M.M.,
Stewart A.M., Cotton M.D., Pratt M.S., Phillips C.A., Richardson D.,
Heidelberg J., Sutton G.G., Fleischmann R.D., Eisen J.A., White O.,
Salzberg S.L., Smith H.O., Venter J.C., Fraser C.M.;
"Evidence for lateral gene transfer between Archaea and Bacteria from
                                                                                                                                                                                                                                                                                                                                                                                      preproparathyroid hormone.";
J. Bone Miner. Res. 3:689-698(1988).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN
BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            30-MAY-2000 (Rel. 39, Last sequence update)
30-MAY-2000 (Rel. 39, Last annotation update)
LEUCYL-TRNA SYNTHETASE (EC 6.1.1.4) (LEUCINE--TRNA LIGASE) (LEURS).
                                                                                        "Nucleotide sequence of the DNA complementary to avian (chicken) preproparathyroid hormone mRNA and the deduced sequence of the
                                                                                                                                                                                                                                                                                                 Khosla S., Demay M., Pines M., Hurwitz S., Potts J.T. Jr., Kronenberg H.M.; "Nucleotide sequence of cloned cDNAs encoding chicken
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      PARATHYROID HORMONE.
B309D8E772997F6E CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        .5e-07;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 90; DB 1;
Pred. No. 9.5e-07
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      824 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 7; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Thermotogales; Thermotoga.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                 MEDLINE-89219100; PubMed-2710135;
                                                                                                                                                                                                                                                                        MEDLINE-89284968; PubMed-3251402;
                                                                                                                                                        hormone precursor.";
Mol. Endocrinol. 3:325-331(1989)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Pfam; PF01279; Parathyroid; 1. PROSITE; PS00335; PARATHYROID;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last seq
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    13943 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     EMBL; M31604; AAA49093.1; -. EEMBL; M36522; AAB02866.1; -. PIR; A34937; A34937. HSSP; P01270; 12WA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          65.7%;
59.3%;
                                                         Russell J., Sherwood L.M.; "Nucleotide sequence of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match 65.77
Best Local Similarity 59.33
Matches 16; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            InterPro; IPR001415; -.
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31
119
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 119 AA;
                                                                                                                                                                                                                                             SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         NCBI_TaxID=2336;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Signal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       LEUS OR TM0168
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      SYL_THEMA
Q9WY15;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Bacteria;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Hormone;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SEQUENCE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CHAIN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SYL_THEMA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT
      RRARRER RRARRE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ö
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                 ribonucleic acid.";
Endocrinology 136:5600-5607(1995).
-!- FUNCTION: PTH ELEVATES CALCIUM LEVEL BY DISSOLVING THE SALTS IN
BONE AND PREVENTING THEIR RENAL EXCRETION.
-!- TISSUE SPECIFICITY: HYPOTHALANUS AND PARATHYROID GLAND.
-!- SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.
                                                                                                                   Schmelzer H.J., Gross G., Mayer H.;
"Nucleotide sequence of cloned cDNA encoding rat prepro parathyroid
                                                                                                                                                                                                                                                                                                                                                           Nutley M.T., Parimi S.A., Harvey S.;
"Sequence analysis of hypothalamic parathyroid hormone messenger
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ó
                                                                                                                                                                                                                                                                  SEQUENCE OF 32-115 FROM N.A. STRÖUENCE OF 32-115 FROM N.A. STRAIN-SPRACHE-DAWLEY; TISSUE-Brain, Parathyroid, and Liver; WEDLINE-9679910; PubMed=7588314; Nutley M.T., Parimi S.A., Harvey S.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            84.7%; Score 116; DB 1; Length 115; 81.5%; Pred. No. 1.2e-10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Indels
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A -> T (IN REF. 3).
V -> I (IN REF. 3).
V -> G (IN REF. 3).
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01-APR-1990 (Rel. 14, Last sequence update)
15-JUL-1998 (Rel. 36, Last annotation update)
PARATHYROID HORMONE PRECURSOR (PTH).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         119 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   3; Mismatches
Nucleic Acids Res. 15:6740-6740(1987)
                                                                                                                                                                                                               Adv. Gene Technol. 21:228-229(1984).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  33 VSEIQLMHNLGKHLASVERMOWLRKKL 59
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    EMBL; X05721; CAA29192.1; -.
EMBL: M54875; AAA57156.1; -.
EMBL: S80127; -; NOT_ANNOTATED_CDS.
PIR; A05091; A05091.
PIR; A26806; A26806.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Pfam; PF01279; Parathyroid; 1.
PROSITE; PS00335; PARATHYROID; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  25
31
115 P.
18 C
23 A
33 V
62 V
12722 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            EMBL; K01268; AAA41979.1; -.
                                                            SEQUENCE OF 10-115 FROM N.A.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     InterPro; IPR001415;
                                                                                        TISSUE=Parathyroid;
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les 22; Conser
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NCBI_TaxID=9031;
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SIGNAL
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P15743;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CONFLICT
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                                                                                                                                                                                   hormone
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RESULT 7
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Best Loc Matches

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                                                                                                                                                                                                                                                                                                                                            This SWISS-PROT entry is copyright. It is produced through a collaboration between the SWISS Institute of Bioinformatics and the EMBL outstation the Buropean Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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STRAIN=ATCC 57210 / B31;
STRAIN=ATCC 57210 / B31;
MEDLINE=98065943; PubMed=9403685;
Fraser C.M., Casjens S., Huang W.M., Sutton G.G., Clayton R.A.,
Lathigra R., White O., Ketchum K.A., Dodson R., Hickey E.K., Gwinn M.,
Dougherty B., Tomb J.-F., Fleischmann R.D., Richardson D.,
Peterson J., Kerlavage A.R., Quackenbush J., Salzberg S., Hanson M.,
van Vugt R., Palmer N., Adams M.D., Gocayne J.D., Weidman J.,
Ulterbock T., Watthey L., McDonald L., Artiach P., Bowman C.,
Garland S., Fulii C., Cotton M.D., Horst K., Roberts K., Hatch B.,
Smith H.O., Venter J.C.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          -i-FUNCTION: CELL WALL FORMATION. CATALYZES THE TRANSFER OF A GLCNAC SUBUNIT ON UNDECAPRENT-PYROPHOSPHORYL-WURNAC-PENTAREPTIDE (LIPID INTERNEDIATE I) TO FORM UNDECAPRENTL-PYROPHOSPHORYL-WURNAC-(PENTAREPTIDE)GLCNAC (LIPID INTERNEDIATE II) (BY SIMILARITY).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       InterPro; IPKUVLOUK,
PPfam; PF00133; TRNA-Synt_1; 1.
PRINTS; PR00985; TRNA-SYNTLEU.
PROSITE; PS00178; AA_TRNA_LIGASE_I; 1.
Aminoacy1-tRNA synthetase; Protein blosynthesis; Ligase; ATP-binding.
                                                                                                                                                                          PYROPHOSPHATE + L-LEUCYL-TRNA(LEU).
-1- SUBCELLULAR LOCATION: CYTOPLASMIC.
-1- SIMILARITY: BELONGS TO CLASS-I AMINOACYL-TRNA SYNTHETASE FAMILY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               30-MAY-2000 (Rel. 39, Last sequence update)
30-MAY-2000 (Rel. 39, Last annotation update)
UDP-N-ACETYLGLUCOSAMINE--N-ACETYLMURAMYL-(PEWTAPEPTIDE)
PYROPHOSPHORYL-UNDECAPRENOL N-ACETYLGLUCOSAMINE TRANSFERASE
(EC 2.4.1.-) (UNDECAPRENYL-PP-MURNAC-PENTAPEPTIDE-UDPGLCNAC GLCNAC TRANSFERASE)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ;
                                                                                                                                            -!- CATALYTIC ACTIVITY: ATP + L-LEUCINE + TRNA(LEU) = AMP +
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 54; DB 1; Length 824; Pred. No. 1.7;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           "Genomic sequence of a Lyme disease spirochaete, Borrelia
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ATP (BY SIMILARITY).
7CB0252A76A844EC CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Borrelia burgdorferi (Lyme disease spirochete).
Bacteria; Spirochaetales; Spirochaetaceae; Borrelia.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               9; Mismatches
                                                           genome sequence of Thermotoga maritima.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           692 MELVNHLSQYLNSVPQEEWNRKLL 715
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5 IQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                EMBL; AE001702; AAD35261.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       95624 MW;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              30-MAY-2000 (Rel. 39, Created)
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ilarity 41.7%;
Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Nature 390:580-586(1997).
                                                                                                              399:323-329(1999).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             InterPro; IPR002300; -. InterPro; IPR002302; -.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
Matches 10; Conserv
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MURG_BORBU
DD 30-MAX DD 30-MAX DD 30-MAX DD 30-MAX DD BE UDP-N-DE UT-DE UT-DE UT-DE UDP-N-DE UT-DE UT-DE UDP-N-DE UT-DE UDP-N-DE UT-DE UT-DE UDP-N-DE UT-DE UT-
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@liber.ch).
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Skelton J., Churcher C.M., Barrell B.G., Rajandream M.A., Walsh S.V., Submitted (JAN-1996) to the EMBL/GenBank/DDBJ databases.
-! - SUBCELLULAR LOCATION: UNCLEAR (PROBBALE).
-! - SIMILARITY: BELONGS TO THE GATA-TYPE TRANSCRIPTION FACTOR FAMILY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Interpro; Irac.
Pfam; PF00320; GATA: 2.
PF10320; GATAZNPINGER.
PROSITE; PS00344; GATAZNPINGER.1; 2.
PROSITE; PS50114; GATA_ZN_FINGER_1; 2.
PROSITE; PS50114; GATA_ZN_FINGER_2; 2.
Transcription regulation; DNA-binding; Zinc-finger; Nuclear protein.
ZN_FING 12 36 GATA_TYPE (BY SIMILARITY).
172 196 GATA_TYPE (BY SIMILARITY).
L -> K (IN REF. 1).
                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
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-!- SUBCELLULAR LOCATION: MEMBRANE ASSOCIATED (BY SIMILARITY).
-!- SIMILARITY: BELONGS TO THE MURG FAMILY.
                                                                                                                                                                                                                                                                  TIGR; BB0767; -.
Transferase; Glycosyltransferase; Cell division; Cell wall;
                                                                                                                                                                                                                                                                                                                                                                                                   Length 363;
                                                                                                                                                                                                                                                                                                                                                                                                                                               Indels
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Hoe K.-L., Park S.-K., Yoo O.-J.J., Yoo H.-S.;
Submitted (MAR-1995) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                              Membrane; Peptidoglycan synthesis.
SEQUENCE 363 AA; 41118 MW; 1BAFA347384DB235 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Schizosaccharomyces pombe (Fission yeast).
Eukaryota; Fungi; Ascomycota; Schizosaccharomycetes;
Schizosaccharomycetales; Schizosaccharomycetaceae;
                                                                                                                                                                                                                                                                                                                                                                                                 38.0%; Score 52; DB 1; 36.0%; Pred. No. 1.5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               01-FEB-1996 (Rel. 33, Last sequence update)
01-NOV-1997 (Rel. 35, Last annotation update)
GAF2 PROTEIN (GAF-2).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 564 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                             7; Mismatches
                                                                                                                                                                                                 or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  217 AEIYFIHQSGKNLNDLSEKNYLRRQ 241
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         3 SEIQLMHNLGKHLNSMXRVEWLRKK 27
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           01-FEB-1996 (Rel. 33, Created)
                                                                                                                                                                                                                                             EMBL; AE001176; AAC67113.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 EMBL; L29051; AAB38022.1; -. EMBL; Z68887; CAA93113.1; -. HSSP; P17429; 5GAT.
                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity 36.v.
Best Local Similarity 36.v.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Schizosaccharomyces.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                GAF2 OR SPAC23E2.01
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            NCBI_TaxID=4896;
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Q10134;
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454 4
508 AA;
                                                                                                                                                                                                                                                                                                                  Escherichia.
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                                                                                                                                                                                                                                                                                                                                                                    STRAIN-K12
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       STRAIN=K12
 CA_BIND
SEQUENCE
                                                   Query Match
                                                                                                                                                               RESULT 12
TORA_ECOLI
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                                                               1;
                                                                                                                                                                                                                                                                                                                                                                                                               SCIENCE 252.951.954(1991).

-!- FUNCTION: MAY PLAY A ROLE IN SIGNAL TRANSDUCTION PATHWAYS THAT INVOLVE CALCIUM. AS A SECOND MESSENGER.

-!- ENZYME REGULATION: ACTIVATED BY CALCIUM. AUTOPHOSPHORYLATION MAY PLAY AN IMPORTANT ROLE IN THE REGULATION OF THE KINASE ACTIVITY.

-!- TISSUE SPECIFICITY: FOUND THROUGHOUT THE PLANT.

-!- MISCELLANBOUS: THERE ARE MULTIPLE CDPK ISOPORAINS IN SOYBEAN.

-!- SIMILARITY: WITH THE CONSERVED CATALYTIC DOMAINS OF SER/THR-PROFEIN KINASES. BELONGS TO THE CAMK SUBFAMILY.

-!- SIMILARITY: CONTAINS 4 EF-HAND CALCIUM-BINDING DOMAINS.
                                                               Gaps
                                                                                                                                                                                                                                                                                                                                                                                           "A calcium-dependent protein kinase with a regulatory domain similar
                                                                                                                                                                                                                                                   Glycine max (Soybean).
Eukaryota; Viridiplantae; Embryophyta; Tracheophyta; Spermatophyta;
Magnoliophyta; eudicotyledons; core eudicots; Rosidae; eurosids I;
Fabales; Fabaceae; Papilionoideae; Glycine.
                                                               5
                                                                                                                                                                                                                                                                                                                                                                  Harper J.F., Sussman M.R., Schaller G.E., Putnam-Evans C.,
Charbonneau H., Harmon A.C.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PROSITE; PS00107; PROTEIN_KINASE_ATP; 1.
PROSITE; PS001108; PROTEIN_KINASE_ST; 1.
PROSITE; PS50011; PROTEIN_KINASE_DOM; 1.
Transferase; Serine/threonine-protein kinase; ATP-binding;
                                     37.2%; Score 51; DB 1; Length 564, 47.6%; Pred. No. 3.3;
                                                                                                                                                                                                01-DEC-1992 (Rel. 24, Created)
01-DEC-1992 (Rel. 24, Last sequence update)
30-MAY-2000 (Rel. 39, Last annotation update)
CALCIUM-DEPENDENT PROTEIN KINASE SK5 (EC 2.7.1.-) (CDPK).
                                                               5; Indels
 0ED74CE0B6E210B7 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PROTEIN KINASE.
ATP (BY SIMILARITY).
ATP (BY SIMILARITY).
BY SIMILARITY.
SITE 1 (POTEWTIAL).
SITE 2 (POTEWTIAL).
SITE 3 (POTEWTIAL).
                                                                                                                                                                           508 AA.
                                                               4; Mismatches
                                                                                                                                                                           PRT;
                                                                                                                                                                                                                                                                                                                                          STRAIN=CV. WILLIAMS;
MEDLINE=91240279; PubMed=1852075;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Phosphorylation.
                                                                                                    519 IQELHNLNQHIQQID--EWLR 537
60611 MW;
                                                                                    5 IQLMHNLGKHLNSMXRVEWLR 25
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         PROSITE; PS00018; EF_HAND; 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       EMBL; M64987; AAB00806.1; -
                                                               10; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Pfam; PF00036; efhand; 4.
Pfam; PF00069; pkinase; 1
                                                                                                                                                                            STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  HSSP; Q63450; 1A06.
InterPro; IPR000719; -.
InterPro; IPR002048; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      InterPro; IPR002290; -
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564 AA;
                                                  Best Local Similarity
                                                                                                                                                                                                                                                                                                                              SEQUENCE FROM N.A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Calcium-binding; DOMAIN 34
NP_BIND 40
BINDING 63
                                                                                                                                                                                                                                                                                                      NCBI_TaxID=3847;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 HSSP; Q63450;
                                                                                                                                                                                                                                                                                                                                                                                                      to calmodulin
                                                                                                                                                                         CDPK_SOYBN P28583;
 SEQUENCE
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CA_BIND
                                      Query Match
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                                                                                                                                                  RESULT 11
CDPK_SOYBN
                                                            Matches
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Ueguchi C., Kakeda M., Yamada H., Mizuno T.;
"An analogue of the DnaJ molecular chaperone in Escherichia coli.";
Proc. Natl. Acad. Sci. U.S.A. 91:1054-1058(1994).
-!- FUNCTION: REDUCES TRIBETHYLAMINE-N-OXIDE (TWAO) INTO
TRIMETHYLAMINE; AN ANAEROBIC REACTION COUPLED TO ENERGY-YIELDING
                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Oshima T., Aiba H., Baba T., Fujita K., Hayashi K., Honjo A.,
Ikemoto K., Inada T., Itoh T., Kajihara M., Kanai K., Kashimoto K.,
Kimura S., Kitagawa M., Makino K., Masuda S., Miki T., Mizobuchi K.,
Mori H., Motomura K., Nakamura Y., Nashimoto H., Nishio Y., Saito N.
Sampel G., Seki Y., Tagami H., Takemoto K., Wada C., Yamamoto Y.,
Yano M., Horiuchi T.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Mejean V., Lobbi-Nivol C., Lepelletier M., Giordano G., Chippaux M.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            "TMAO anaerobic respiration in Escherichia coli: involvement of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                [2]
SEQUENCE FROM N.A.
SEQUENCE FROM N.A.
MEDLINE-97426617; PubMed=9278503;
Blattner F.R., Plunkett G. III, Bloch C.A., Perna N.T., Burland V., Riley M., Collado-Vides J., Glasner J.D., Rode C.K., Mayhew G.F., Greegor J., Davis N.W., Kirkpatrick H.A., Goeden M.A., Rose D.J., Mau B., Shao Y.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               -!- CATALYTIC ACTIVITY: NADH + TRIMETHYLAMINE-N-OXIDE = NAD(+) + TRIMETHYLAMINE + H(2)0.
-! TRIMETHYLAMINE + H(2)0.
-! SUBCELLULAR LOCATION: PERIPLASMIC.
-! SIMILARITY: BELONGS TO THE PROKARYOTIC MOLYBDOPTERIN-CONTAINING
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Yano M., Horiuchi T.;
"A 718-kb DNA sequence of the Escherichia coli K-12 genome
corresponding to the 12.7-28.0 min region on the linkage map.";
                                                                                                                                                                                                   Ξ,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    01-FEB-1994 (Rel. 28, Created)
01-NOV-1997 (Rel. 35, Last sequence update)
15-JUL-1999 (Rel. 38, Last annotation update)
TRIMETHYLAMINE-N-OXIDE REDUCTASE PRECURSOR (EC 1.6.6.9) (TMAO
                                                                                                                              Length 508;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 "The complete genome sequence of Escherichia coli K-12."; Science 277:1453-1474(1997).
                                                                                                                                                                                                   Indels
165 SITE 4 (POTENTIAL).
57169 WW; AFCEDC51224192E4 CRC64;
                                                                                                                                  ;;
                                                                                                                                  DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      848 AA.
                                                                                                                              Score 50.5; Dl
Pred. No. 3.5;
5; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE FROM N.A., AND SEQUENCE OF 40-46.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      REDUCTASE) (TRIMETHYLAMINE OXIDASE).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Mol. Microbiol. 11:1169-1179(1994)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            MEDLINE=94293785; PubMed=8022286;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  MEDLINE=97061202; PubMed=8905232;
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                                                                                                                              36.9%;
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EIQIMHHLSEHAN-VVRIE 99
                                                                                                                                                                                                                                                                    4 EIQLMHNLGKHLNSMXRVE 22
                                                                                                                                                                 Best Local Similarity 52.6
Matches 10; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA Res. 3:137-155(1996).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         STANDARD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Escherichia coli
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TORA_ECOLI
P33225; P78227;
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ACT_SITE
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                          This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation-the Buropean Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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01-NOV-1988 (Rel. 09, Last sequence update)
01-NOV-1997 (Rel. 35, Last annotation update)
BETA PLATELET-DERIVED GROWTH FACTOR RECEPTOR PRECURSOR (EC 2.7.1.112)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Mus musculus (Muse).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
NGBI_TaxID=10090;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          7; Indels
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                                                                                                                                                                                                                                                                                                                                                                                           Plam; PFU0364; molybdopterin; 1.
PROSITE; PS00551; MOLYBDOPTERIN_PROK_1; FALSE_NEG.
PROSITE; PS00490; MOLYBDOPTERIN_PROK_2; 1.
PROSITE; PS00932; MOLYBDOPTERIN_PROK_3; 1.
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G (IN REF. 1).

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P -> L (IN REF. 1
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00 -> HE (
                                                                                                                                                                                                                     EMBL; AE000201; AAC74082.1; -.
EMBL; D90736; BAA35139.1; -.
EMBL; D90737; BAA35764.1; -.
EMBL; D16500; -; NOT_ANNOTATED_CDS.
BPTR; S34222; S34222.
HSSP; Q57366; 1CXT.
                                                                                                                                                                                                                                                                                                                                             Pfam; PF01568; Molydop_binding; 1. Pfam; PF00384; molydop_terin; 1.
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MEDLINE-87014762; PubMed=3020426;
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                        OXIDOREDUCTASE FAMILY
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hes 9; Conserv
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PDGFRB OR PDGFR.
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PGDR_MOUSE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ö
                                                                                                  -i- SUBCELLULAR LOCATION: TYPE I MEMBRANE PROTEIN.
-i- SIMILARITY: BELONGS TO THE CSF-1/PDGF RECEPTOR FAMILY OF TYROSINE-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
Nature 323:226-232(1986).
-!- FUNCTION: THIS RECEPTOR BINDS PLATELET-DERIVED GROWTH FACTOR AND
HAS A TYROSINE-PROTEIN KINASE ACTIVITY. THIS RECEPTOR BINDS
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                                                                   SUBUNIT: DIMER OF EITHER ALPHA-ALPHA, BETA-BETA OR ALPHA-BETA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Tyrosine-protein kinase; Receptor; Transmembrane; Glycoprotein;
Transferase; Phosphorylation; ATP-binding; Immunoglobulin domain;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PLATELET - DERIVED GROWTH FACTOR
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Pred. No. 26;
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                                                                                                                                   PROTEIN KINASES.
SIMILARITY: CONTAINS 5 IMMUNOGLOBULIN-LIKE DOMAINS.
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IG-LIKE C2-TYPE DOMAIN.
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InterPro; IPR00306; -
Pfam: PF00040; ig; 4.
Prom: PF00069; pkinase; 2.
PROSITE; PS00109; PROTEIN_KINASE_ATP; 1.
PROSITE; PS00109; PROTEIN_KINASE_ATP; 1.
PROSITE; PS00109; PROTEIN_KINASE_ATP; 1.
PROSITE; PS00240; RECEPFOR TYR KIN III; 1.
PROSITE; PS50011; PROTEIN_KINASE_DOM; 1.
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53.3%;
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                                                    SPECIFICALLY TO PDGF-B
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InterPro; IPR000719; -.
InterPro; IPR001245; -.
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Best Local Similarity
Matches 8; Conserv
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DISEASE: INVOLVED IN A FORM OF CHRONIC MYELDMONOCYTIC LEUKEMIA (CMML), CHARACTERIZED BY ABNOMAL CLONAL MYELDMONOCYTIC LEUKEMIA AND BY PROGRESSION TO ACUTE MYELOGENOUS LEUKEMIA (AML). IT IS BY PROGRESSION TO ACUTE MYELOGENOUS LEUKEMIA (AML). IT IS BETWERN THE ETS-LIKE PROTEIN TEL AND PEDF RECEPTOR B (PDGFRB). SIMILARITY: BELONGS TO THE CSF-1/PDGF RECEPTOR B (PDGFRB).
                                                                                                                                01-WAR-1989 (Rel. 10, Last sequence update)
15-JUL-1998 (Rel. 36, Last annotation update)
BETA PLATELET-DERIVED GROWTH FACTOR RECEPTOR PRECURSOR (EC 2.7.1.112)
(PDGF-R-BETA) (CD140B ANTIGEN).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Kazlauskas A., Cooper J.A.;
"Autophosphorylation of the PDGF receptor in the kinase insert region regulates interactions with cell proteins.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           FUNCTION: THIS RECEPTOR BINDS PLATELET-DERIVED GROWTH FACTOR AND HAS A TYROSINE-PROTEIN KINASE ACTIVITY, THIS RECEPTOR BINDS SPECIFICALLY TO PDGF-B.
                                                                                                                                                                                                                                                                                  MEDIINE-88217915; PubMed-2835772; Gronwald R.G.K., Grant F.J., Haldeman B.A., Hart C.E., O'Hara P.J., Hagen F.S., Ross R., Bowen-Pope D.F., Murray M.J.; "Cloning and expression of a cDNA coding for the human platelet-derived growth factor receptor: evidence for more than one
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Roberts W.M., Look A.T., Roussel M.F., Sherr C.J.; "Tandem linkage of human CSF-1 receptor (c-fms) and PDGF receptor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SUBUNIT: DIMER OF EITHER ALPHA-ALPHA, BETA-BETA OR ALPHA-BETA
                                                                                                                                                                                                                Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                     Claesson-Welsh L., Eriksson A., Moren A., Severinsson L., Ek B., Oestman A., Betsholtz C., Heldin C.-H.; "cDNA cloning and expression of a human platelet-derived growth factor (PDGF) receptor specific for B-chain-containing PDGF molecules.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Chi K.D., McPhee R.A., Wagner A.S., Dietz J.J., Pantazis Goustin A.S.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Submitted (MAY-1997) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SIMILARITY: CONTAINS 5 IMMUNOGLOBULIN-LIKE DOMAINS
                                                                                                                                                                                                                                                                                                                                                                   Proc. Natl. Acad. Sci. U.S.A. 85:3435-3439(1988)
                                                                                           1106 AA
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                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A. MEDLINE-89096941; Pubmed=2850496;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AUTOPHOSPHORYLATION SITES.
MEDLINE=89376563; PubMed=2550144;
                                                                                                                       Created)
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648 MSELKIMSHLGPHLN 662
                                                                                           STANDARD;
2 VSEIQLMHNLGKHLN 16
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Cell 55:655-661(1988).
                                                                                                                                                                                                       Homo sapiens (Human).
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                                                                                                                                                                                                                                            NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                        receptor class.
                                                                                         PGDR_HUMAN
P09619;
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the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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PROSITE; PS00109; PROTEIN_KINASE_TYR; 1.
PROSITE; PS00101; PROTEIN_KINASE_LOM; 1.
PROSITE; PS00240; RECEPTOR_TYKIN_III; 1.
Tyrosine-protein kinase; Receptor; Transmembrane; Glycoprotein; Transferase; Phosphorylation; ATP-binding; Immunoglobulin domain;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Mus musculus (Mouse).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
NCBI_TaxID=10090;
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O1-OCT-2000 (Rel. 40, Last annotation update)
PARATHYROID HORMONE-RELATED PROTEIN PRECURSOR (PTH-RP) (PLP)
PATHLH OR PTHRP
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PHOSPHORYLATION (AUTO-).
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                                                                                                           EMBL; J03278; AAA60049.1; -. EMBL; M21616; AAA36427.1; -. EMBL; U33172; AAC51675.1; -.
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Pfam; PF00069; pkinase; 2.
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InterPro; IPR001245; -.
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InterPro; IPR003006; -.
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RA MANGIN M. Ikeda K., Broadus A.E.;

"Structure of the mouse gene encoding parathyroid hormone-related
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C -1 - SIMILARITY: BELONGS TO THE PARATHYROID HORMONE FAMILY.

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C ENE ENDOSEN BIOINFORMATICS INSTITUTE. There are no restrictions on its
C c case by non-profit institutions as its content is in no way
C modified and this statement is not removed usage by and for commercial
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EMBL: M60058; AAA6359.1;
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C Thirder Post IPROULE THE PARATHYROID HORMONE-RELATED PROTEIN.

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Search completed: August 31, 2001, 16:01:46 Job time: 110 sec

Gaps

6; Indels

33.6%; Score 46; DB 1; Length 175; 47.4%; Pred. No. 5.5;

4; Mismatches

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Query Match
Best Local Similarity 47.4'
Matches 9; Conservative

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Search time 12.82 Seconds (without alignments) 166.372 Million cell updates/sec
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Copyright (c) 1993 - 2000 Compugen Ltd.
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1 XVSEIQLMHNLGKHLNSMXRVEWLRKKL 28 US-09-448-867-1 Perfect score: Sequence: Title:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

219241 seqs, 76174552 residues

219241 Total number of hits satisfying chosen parameters:

length: 0 length: 2000000000 sed Minimum DB Maximum DB

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

PIR_68:*
1: pir1:*
2: pir2:*
3: pir3:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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A; Molecule type: DNA A; Residues: 1-115 <VAS>

A;Cross-references: GB:J00301; NID:g190702; PIDN:AAA60215.1; PID:g190704
B;Yamaguchi, T.; Fukase, M.; Suginoto, T.; Kido, H.; Chihara, K.
Biol. Chem. Hoppe-Seyler 375, 821-824, 1994
A;Title: Purification of meprin from human kidney and its role in parathyroid hormone A;Reference number: S53790; MUID:95225988
A;Accession: S53790

A; Molecule type: protein
A; Residues: 'X', 33, 'X', 35, 46; 65-84; 105-110 < YAM>
A; Residues: 'X', 33, 'X', 35, 46; 65-84; 105-110 < YAM>
A; Note: peptides generated in vitro and in vivo by meprin; peptide cleavage also occu
R; Jacobs, J.W.; Kemper, B.; Niall, H.D.; Habener, J.F.; Potts Jr., J.T.
Nature 249, 155-157, 1974
A; Title: Structural analysis of human proparathyroid hormone by a new microsequencing
A; Reference number: A93169; MUID: 74174967
A; Accession: A93169

A; Wolecule type: protein A; Residues: 26.37 <JAC> A; Residues: 26.37 <JAC> Colstad, O.K.; Reppe, S.: Gabrielsen, O.S.; Hartmanis, M.; Blingsmo, O.R.; Gautvik, Eur. J. Biochem. 205, 311-319, 1992 A; Title: Isolation and characterization of two biologically active O-glycosylated for ation.

A; Reference number: S21199; MUID: 92209518 A; Accession: S21199

A; Molecule type: protein
A; Residues: 32-114,'N' <0LS>
A; Residues: 32-114,'N' <0LS>
A; Residues: 23-114,'N' <0LS>
A; Note: cloned sequence expressed in Saccharomyces cerevisiae exhibited O-glycosylati
B; Niall, H.D.; Sauer, R.T.; Jacobs, J.W.; Keutmann, H.T.; Segre, G.V.; O'Riordan, J.L.
Proc. Natl. Acad. Sci. U.S.A. 71, 384-388, 1974
A; Title: The amino-acid sequence of the amino-terminal 37 residues of human parathyro
A; Reference number: A93789; MUID:74111656
A; Accession: A93789

A; Molecule type: protein A; Residues: 32-68 <NIA> R; Brewer Jr., H.B.; Fairwell, T.; Ronan, R.; Sizemore, G.W.; Arnaud, C.D. Proc. Natl. Acad. Sci. U.S.A. 69, 3585-3588, 1972 A; Title: Human parathyroid hormone: amino-acid sequence of the amino-terminal residue A; Reference number: A93783; MUID:73070429

A; Molecule type: protein

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A;Cross-references: GB:K01938
R;Kronenberg, H.M.; McDevitt, B.E.; Majzoub, J.A.; Nathans, J.; Sharp, P.A.; Potts Jr Proc. Natl. Acad. Sci. U.S.A. 76, 4981-4985, 1979
A;Title: Cloning and nucleotide sequence of DNA coding for bovine preproparathyroid h A;Reference number: A93835; MUID:80056617
A;Accession: A93835
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 A Accession: A90376
A, Molecule type: protein
A, Residues: 32-109 cSAU>
R; Brewer Jr., H.B.; Fairwell, T.; Rittel, W.; Littledike, T.; Arnaud, C.D.
Am. J. Med. 56, 759-766, 1974
A; Title: Recent studies on the chemistry of human, bovine and porcine parathyroid hor A; Reference number: A90030; MUID:74173303
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parathyroid hormone precursor - pig
C;Species: Sus scrofa domestica (domestic pig)
C;Date: 24-Apr-1984 #sequence_revision 12-Apr-1996 #text_change 18-Jun-1999
C;Accession: Bz6806; A90370; A90376; A01535
R;Schmelzer, H.J.; Gross, G.; Widera, G.; Mayer, H.
Wucleic Acids Res. 15, 6740, 1987
A;Title: Nucleotide sequence of a full-length cDNA clone encoding preproparathyroid A;Reference number: Az6806; MUID:87316938
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A;Note: the authors translated the codon GAA for residue 50 as Gly
R;Hamilton, J.W.; Niall, H.D.; Jacobs, J.W.; Keutmann, H.T.; Potts Jr., J.T.; Cohn,
                                                                                                                                                                                                                                                                                                                                                                                                        A;Cross-references: GB:X05722; GB:Y00409; NID:91838; PIDN:CAA29193.1; PID:91839
R;Chu, L.L.H.; Huang, W.Y.; Littledike, E.T.; Hamilton, J.W.; Cohn, D.V.
Biochemistry 14, 3631-3635, 1975
A;Title: Porcine proparathyroid hormone. Identification, biosynthesis, and partial A;Reference number: A90390; MUID:76018954
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Hogan, M.L.; Keutmann, H.T.; O'Riordan, J.L.H.; Potts
1974
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C; Date: 23-Oct.1981 #sequence_revision 23-Oct.1981 #text_change 18-Jun-1999
C; Accession: A24949; A93835; A93793; A91648; A93773; I45975; I45976; A01534
R; Weaver, C.A.; Gordon, D.F.; Kissil, M.S.; Mead, D.A.; Kemper, B.
Gene 28, 319-329, 1984
A; Title: Isolation and complete nucleotide sequence of the gene for bovine p
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A; Residues: 26-115 <CHU>
S: Sauer, R.T.; Niall, H.D.; Hogan, M.L.; Keutmann, H.T.; O'Riord
Biochemistry 13, 1994-1999, 1974
A; Title: The amino acid sequence of porcine parathyroid hormone.
A; Reference number: A90376; MUID:74253317
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F;30-64/Domain: parathyroid hormone homology <PTH>
F;32-115/Product: parathyroid hormone #status experimental <MAT>
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Pred. No. 1.1e-11;
                                                                                                                                                                                                                                                                                                               A;Status: not compared with conceptual translation A;Molecule type: mRNA
A;Residues: 1-115 <SÇH>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              C; Superfamily: parathyroid hormone; parathyroid C; Keywords: calcium; hormone; parathyroid gland
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2; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      parathyroid hormone precursor - bovine
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88.9%;
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Matches 24; Conservative
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A; Residues: 1-115 <WEA>
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RESULT PTPG

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A; Molecule type: protein
A; Mesidues: 75-100 (KEZ)
A; Residues: 75-100 (KEZ)
A; Residues: 75-100 (KEZ)
B; Tregear, G.W.; van Rietschoten, J.; Green, E.; Niall, H.D.; Keutmann, H.T.; Parsons, J. Hoppe-Seyler's 2. Physiol. Chem. 355, 415-421, 1974
A; Title: Solid-phase synthesis of the biologically active N-terminal 1-34 peptide of hum A; Reference number: A91660; MUID:75059220
A; Reference number: A91660; MUID:75059220
A; Contents: annotation; synthesis of residues 32-65
A; Note: the biologically active amino-terminal 34 residues of parathyroid hormone were sat renal adenylate cyclase assay and with the bovine hormone's active region in the chic Randerstar, R.H.; Hartmann, A.; Joehl, A.; Kamber, B.; Maier, R.; Riniker, B.; Rittel, Helv. Chim. Acta 56, 470-473, 1973
A; Title: Synthese der Sequenz 1-34 von menschlichem Parat-hormon.
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Proc. Natl. Acad. Sci. U.S.A. 78, 7365-7369, 1981
A; Title: Nucleotide sequence of cloned cDNAs encoding human preproparathyroid hormone.
A; Reference number: 138342; MUID:82150870
                                             and mass spectroscopic identification obtts \mathrm{Jr.}, \mathrm{J.F.}
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    P.M.; Hendy, G.N.; O'Riordan, J.L.
M., and Parsons, J.A., eds., pp.9-
                                                                                                                                                             A;Title: A reinvestigation of the amino-terminal sequence of human parathyroid hormone. A;Reference number: A90387; MUID:75146516
A;Accession: A90387
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A;Residudes: 52-75 <KRS.
R;Keutmann, H.T.; Sauer, M.M.; Hendy, G.N.; O'Riordan, J.L.H.; Potts Jr., J.T.
Biochemistry 17, 5723-5729, 1978
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                                                                                                                                                                                                                                                                                                                                                                                                                                             A; Title: Complete amino acid sequence of human parathyroid hormone.
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A;Residues: 32-52,'0',54-58,'K',60,'L',62-65 <BRE>A;Note: this sequence was determined by sequenator and mass s R:Keutmann, H.T.; Niall, H.D.; O'Riordan, J.L.H.; Potts Jr., Biochemistry 14, 1842-1847, 1975
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R;Keutmann, H.T.; Niall, H.D.; Jacobs, J.W.; Barling,
in Calcium-regulating Hormones, Talmadge, R.V., Owen,
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Pred. No. 7.1e-13;
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0; Mismatches
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A;Cross-references: GDB:119522; OMIM:168450
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A; Accession: A90426
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A;Status: translated from GB/EMBL/DDBJ
A;Molecule type: mRNA
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ilarity 96.3%;
Conservative
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Best Local Similarity
Labes 26; Conserv?
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A; Residues: 61-106,'D',10
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A; Accession: A91648

A; Accession: A93773

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C;Accession: JC4202
R;Rosol, T.J.; Steinmeyer, C.L.; McCauley, L.K.; Groene, A.; DeWille, J.W.; Capen, C. Gene 160, 241-243, 1995
A;Fitle: Sequences of the cDNAs encoding canine parathyroid hormone-related protein a A;Reference number: JC4201; MUID:95369696
A;Accession: JC4202
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C;Species: Rattus norvegicus (Norway rat)
C;Accession: 151851
R;Schmelzer, H.
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C;Species: Rattus norvegicus (Norway rat)
C;Date: 05-Jun 1987 #sequence_revision 05-Jun-1987 #text_change 16-Jul-1999
C;Accession: A05091; A2600
J; Biol. Chem. 259, 3320-3329, 1984
A;Reference number: A05091; MUID:84135846
    C;Species: Canis lupus familiaris (dog)
C;Date: 10-Sep-1995 #sequence_revision 27-Oct-1995 #text_change 16-Jul-1999
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C;Superfamily: parathyroid hormone; parathyroid hormone homology
F;30-64/Domain: parathyroid hormone homology <PTH>
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81.5%; Pred. No. 3.4e-10;
Live 3; Mismatches 2; Indels
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Pred. No. 1.5e-11;
1; Mismatches 2; Indels
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Nucleic Acids Res. 15, 6740, 1987
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88.9%;
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Best Local Similarity 88.9°
Matches 24; Conservative
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Matches 22; Conservative
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A; Residues: 1-115 <ROS>
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Akolecule type: protein
Akolecule type: protein
Akolecule type: 32-115 cares
Akolecule type: 32-115 cares
Akolecule type: 32-115 cares
Akolecule type: 32-115 cares
Akolecule type: brod to a biologically active N-terminal tetratriacontapeptide of parathyr
Akolecule synthesis of a biologically active N-terminal tetratriacontapeptide of parathyr
Akolecule annotation; synthesis of residues 32-65
Akolecule annotation; synthesis of residues 32-65
Akolecule the synthetic peptide was active in vivo and in viro
Akolecule the synthetic peptide was active in vivo and in viro
Akolecule the synthetic peptide was active in vivo and in viro
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Akolecule the synthetic peptide was active in vivo and active in vivo and 
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Proc. Natl. Acad. Sci. U.S.A. 78, 4073-4077, 1981
Affitle: introduction by molecular cloning of artifactual inverted sequences at the A; Reference number: 145975; MUID: 82037778
A; Reference number: 145975; MID: 2007778
A; Recession: 145975
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Proc. Natl. Acad. Sci. U.S.A. 71, 653-656, 1974
A;Title: The N-terminal amino-acid sequence of bovine proparathyroid hormone.
A;Reference number: A93793; MUID:74142666
A;Accession: A93793
A;Molecule type: protein
A;Residues: 26-115 <HAM>
R;Niall, H.D.; Keutmann, H.T.; Sauer, R.; Hogan, M.L.; Dawson, B.F.; Aurbach, Hoppe-Seyler's Z. Physiol. Chem. 351, 1586-1588, 1970
A;Title: The amino acid sequence of bovine parathyroid hormone I.
A;Reference number: A91648; MUID:71076162
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Mol. Cell. Endocrinol. 28, 411-424, 1982
A;Title: Nucleotide sequence of bovine parathyroid hormone messenger RNA. A;Reference number: I45976; MUID:83105964
A;Accession: I45976
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F;26-115/Product: proparathyroid hormone #status experimental <PMAT>
F;26-31/Domain: propeptide #status experimental <PRO>
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C;Reywords: hormone
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A;Residues: 32-115 <NIA>
R;Brewer Jr., H.B.; Ronan, R.
Proc. Natl. Acad. Sci. U.S.A. 67, 1862-1869, 1970
A;Title: Bovine parathyroid hormone: amino acid sequence. A;Reference number: A93773; MUID:71063634
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A; Molecule type: mRNA A; Residues: 1-115 <WE2>

A; Contents: annotation

A; Molecule type: mRNA A; Residues: 1-115 <WE3>

A; Introns: 29/2

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91.2%; 88.9%;

Best Local Similarity 88.9 Matches 24; Conservative

Query Match

parathyroid hormone precursor - dog

ŘESULT JC4202

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A;Cross-references: GB:AE001176; GB:AE000783; NID:g2688699; PIDN:AAC67113.1; PID:g268
A;Experimental source: strain B31
C;Superfamily: murG protein
                                                               A;Molecule type: DNA
A;Residues: 1-824 <ARN>
A;Cross-references: GB:AE001702; GB:AE000512; NID:g4980662; PIDN:AAD35261.1; PID:g498
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      UDP-N-acetylglucosamine--N-acetylmuramyl-(pentapeptide) pyrophosphoryl-undecaprenol
C;Species: Borrelia burgdorferi (Lyme disease spirochete)
C;Date: 13-Feb-1998 #sequence_revision 13-Feb-1998 #text_change 24-Nov-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      C;Accession: F70195
R;Fraser, C.M.; Casjens, S.; Huang, W.M.; Sutton, G.G.; Clayton, R.; Lathigra, R.; son, D.; Peterson, J.; Kerlavage, A.R.; Quackenbush, J.; Salzberg, S.; Hanson, M.; Bowman, C.; Garland, S.; Fujii, C.; Cotton, M.D.; Horst, K.; Roberts, K.; Hatch, Nature 390, 580-586, 1997
A;Authors: Smith, H.O.; Venter, J.C.
A;Title: Genomic sequence of a Lyme disease spirochaete, Borrelia burgdorferi.
A;Reference number: A70100; MUID:98065943
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A;Status: preliminary; nucleic acid sequence not shown; translation not shown A;Molecule type: DNA A;Residues: 1-363 <KLE>
                                                                                                                                                                                                                                      A;Gene: TW0168
C;Superfamily: leucine--tRNA ligase
C;Keywords: aminoacyl-tRNA synthetase; ligase; protein biosynthesis
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Pred. No.
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No.
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                                                                                                                                                                      A; Experimental source: strain MSB8
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36.0%;
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Best Local Similarity
Matches 10; Conserv
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                                      A;Status: preliminary
   A; Accession: F72408
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A34937
parathyroid hormone precursor - chicken
C; Species: Gallus gallus (chicken)
C; Date: 07-Sep-1990 #sequence_revision 07-Sep-1990 #text_change 16-Jul-1999
C; Accession: A34937; 150411
R; Russell, J.; Sherwood, L.M.
Mol. Endocrinol. 3, 325-331, 1989
A;Title: Nucleotide sequence of the DNA complementary to avian (chicken) preproparathyro
A; Reference number: A34937; MulD:89219100
A; Reference number: A34937
A; Molecule type: mRNA
A; Residues: 1-119 < RUS;
A; Molecule type: mRNA
A; Residues: 1-119 < RUS;
A; Molecule type: mRNA
A; Residues: 1-119 < RUS;
A; Molecule type: mRNA
A; Residues: 1-119 < RUS;
A; Molecule type: mRNA
A; Residues: 1-110 < RUS;
A; Molecule type: mRNA
A; Residues: 1-110 < RUS;
A; Molecule type: mRNA
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A; Molecule type: mRNA
A; Residues: 1-110 < RUS;
A; Molecule type: mRNA
A; Residues: 1-110 < RUS;
A; Molecule type: mRNA
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A; Molecule type: mRNA
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A; Molecule type: mRNA
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A; Molecule type: mRNA
A; Residues: 1-10 < RUS;
A; Molecule type: mRNA
A; Residues: 1-10 < RUS;
A; Molecule type: mRNA
A; Russello type: 
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Garrett, M.M.; Stewart, A.M.; Cotton, M.D.; Pratt, M.S.; Phillips, C.A.; Richardson, D.;
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A;Reference number: A72200; WUID:99287316
A;Title: Nucleotide sequence of cloned cDNA encoding rat prepro parathyroid hormone.
A;Reference number: 151851
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A; Residues: 1-19 < KHO>
A; Residues: 1-19 < KHO>
A; Cross-references: GB-M36522; NID:q212591; PIDN:AAB02866.1; PID:q212592
C; Superfamily: parathyroid hormone; parathyroid hormone homology
F; 1-25/Domain: signal sequence #status predicted <SIG>
F; 26-31/Domain: propeptide #status predicted <PRO>
F; 30-64/Domain: parathyroid hormone homology <PTH>
F; 32-119/Product: parathyroid hormone #status predicted <MAT>
                                                                                                                               A;Molecule type: mRNA
A;Residues: 1-105 <RES>
A;Cross-references: GB:M54875; NID:g601932; PIDN:AAA57156.1; PID:g601933
C;Genetics:
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C;Superfamily: parathyroid hormone; parathyroid hormone homology
F;20-54/Domain: parathyroid hormone homology <PTH>
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4.3e-10;
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                                                                                                   A;Status: preliminary; translated from GB/EMBL/DDBJ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 90; DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                4; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          7; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 115;
Pred. No. 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          23 ISEIQLMHNLGKHLASVERMQWLRKKL 49
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2 VSEIQLMHNLGKHLNSMXRVEWLRKKL 28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       65.7%;
59.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                            83.9%;
77.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                21; Conservative
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Matches 16; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
                                                                      A; Accession: I51851
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family protein Gaf2p – fission yeast (Schizosaccharomy pombe
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A;Experimental source: strain 972h-; cosmid c23E2
       GATA-type transcription factor family protein Gaf2p - fission yeast (Schizos C;Species: Schizosaccharomyces pombe
C:Date: 03-Dec-1999 #sequence_revision 03-Dec-1999 #text_change 03-Dec-1999
C;Accession: T38291
                                                                                                                                              R;Skelton, J.; Churcher, C.M.; Barrell, B.G.; Rajandream, M.A.; Walsh, submitted to the EMBL Data Library, January 1996
A;Reference number: 221784
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 564;
                                                                                                                                                                                                                                                                                                        A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
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A;Status: preliminary
A;Molecule type: DNA
A;Rosidues: 1-336 <STO>
A;Cross-references: GB:AP001518; GB:BA000004; NID:g10175792; PIDN:BAB06973.1; GSPDB:G
A;Experimental source: strain C-125
C;Genetics:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     A;Cross-references: GB.SE00201; GB:U00096; NID:92367113; PIDN:AAC74082.1; PID:917872
A;Experimental source: strain K-12, substrain MG1655
R;Mejean, V; Tobbi-Nivol, C.; Lepelletier, M.; Giordano, G.; Chippaux, M.; Pascal, M
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       A;Residues: 1-172,'R',174-175,'R',177-255,'R',257,'S',259-280,'G',283-324,'E',326-347
A;Cross-references: EMBL:X73888; NID:9556701; PIDN:CAA52095.1; PID:9806336
A;Experimental source: strain K-12
hypothetical protein BH3254 [imported] - Bacillus halodurans (strain C-125) C.Species: Bacillus halodurans C.Date: 01-Dec-2000 #sequence_revision 01-Dec-2000 #text_change 31-Dec-2000 C.Accession: F84056 R.Takami, H.; Nakasone, K.; Takaki, Y.; Maeno, G.; Sasaki, R.; Masui, N.; Fuji, F.; Nucleic Acids Res. 28, 4317-4331, 2000 A.Title: Complete genome sequence of the alkaliphilic bacterium Bacillus halodurans A; Reference number: A83650; MUID:20263314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       trimethylamine-N-oxide reductase (EC 1.6.6.9) precursor - Escherichia coli C; Species: Escherichia coli C; Species: Escherichia coli C; Date: 12-Sep-1997 #sequence_revision 17-Sep-1997 #text_change 24-Sep-1999 C; Accession: C64841; S43698; S34222 C; Accession: C64841; S43698; S34222 C; Accession: C64841; S43698; S34222 C; Accession: C64841; Shunkett III, G.; Bloch, C.A.; Perna, N.T.; Burland, V.; Riley, A; Bilattner, F.R.; Plunkett III, G.; Bloch, C.A.; Perna, N.T.; Burland, V.; Riley, A; Rose, D.J.; Mau, B.; Shao, Y. Science 277, 1453-1462, 1997 A; Fiterence number: A64720; MuID:97426617 A; Reference number: A64720; MuID:97426617
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A;Title: TMAO anaerobic respiration in Escherichia coli: involvement of the A;Reference number: S43697; MUID:94293785
A;Accession: S43698
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       A) Description: reduces trimethylamine-N-oxide into trimethylamine C; Superfamily: trimethylamine-N-oxide reductase C; Keywords: molybdenum; NAD; oxidoreductase; periplasmic space F; F: 1-39/Domain: signal sequence #status predicted <SIG>F; 40-848/Product: trimethylamine-N-oxide reductase #status predicted
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 336,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 848
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Indels
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19;
                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    4; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 50;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 50;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             298 VLAILLHFIGPALISLSLSEWFRKR 323
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            2 VSEIQLMHNLGKHLNSMXRVEWLRKK 27
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           36.5%;
45.0%;
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42.3%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11; Conservative
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               A; Molecule type: DNA
                                                                                                                                                                                                                                                                                                                                                                                A:Gene: BH3254
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
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C; Species: Schizosaccharomyces pomber (C; Date: 11-Jan-2000 #text_change 21-Jul-2000 C; Accession: T43298
R; Hoe, K.L.; Won, M.S.; Yoo, O.J.; Yoo, H.S.
Biochem. Mol. Biol. Int. 39, 127-135, 1996
A; Title: Molecular cloning of GAF2, a Schizosaccharomyces pombe GATA factor, which has A; Reference number: Z22402; MUID:96392547
A; Accession: T43298
A; Status: preliminary; translated from GB/EMBL/DDBJ
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                                                       Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          A;Gene: gaf2
C;Keywords: transcription factor; transcription regulation; zinc finger
                                                                                                                                                                                                                                    T43298 transcription factor gaf2 - fission yeast (Schizosaccharomyces pombe)
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Pred. No. 9.4;
5; Mismatches 3;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      37.2%; Score 51; DB 47.6%; Pred. No. 8.8;
                          Pred. No. 8.8;
                                                    Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Mismatches
                                                  4;
                                                                                                                               519 IQELHNLNQHIQQID--EWLR 537
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                                                                                                5 IQLMHNLGKHLNSMXRVEWLR 25
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52.6%;
                            47.68;
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Best Local Similarity 52.6
Matches 10; Conservative
                                                  Conservative
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                          Best Local Similarity
Matches 10; Conserv
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nes 10; Conserv
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A; Residues: C;Genetics:

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trimethylamine N-oxide reductase subunit [imported] - Escherichia coli (strain 0157:H7) C; Species: Escherichia coli (C; Date: 16-Feb-2001 #sequence_revision 16-Feb-2001 #text_change 31-Mar-2001 (C; Date: 16: Feb-2001 #sequence_revision 16-Feb-2001 #text_change 31-Mar-2001 R; Perna, N.T.; Plunkett III, G.; Burland, V.; Mau, B.; Glasner, J.D.; Rose, D.J.; Mayhev iller, L.; Grotbeck, E.J.; Davis, N.W.; Lim, A.; Dimalanta, E.; Potamousis, K.; Apodaca, Nature 409, 529-533, 2001 A; Fille: Genome sequence of enterohemorrhagic Escherichia coli 0157:H7.
A; Reference number: A85480; MUID:21074935; PMID:11206551
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A;Status: preliminary
A;Molecule type: DNA
A;Molecule type: DNA
A;Residues: 1-848 <STO>
A;Cross-references: GB:AE005174; NID:g12514264; PIDN:AAG55544.1; GSPDB:GN00145; UWGP:214
A;Experimental source: strain 0157:H7, substrain EDL933
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       A, Gene: torA C; Superfamily: trimethylamine-N-oxide reductase
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Search completed: August 31, 2001, 16:01:01 Job time: 65 sec

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8 MHNLGKHLNSMXRVEWLRKK 27

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0; Gaps

Score 50; DB 2; Length 848; Pred. No. 19; 4; Mismatches 7; Indels

Ouery Match 36.5%; Best Local Similarity 45.0%; Matches 9; Conservative